PATENT SPECIFICATION

(11) 1 513 672

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(21) Application No. 20686/75

(22) Filed 15 May 1975

(31) Convention Application No. 70096

(32) Filed 16 May 1974

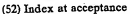
(31) Convention Application No. 71849

(32) Filed 14 Feb. 1975 in

(33) Luxembourg (LU)

(44) Complete Specification published 7 June 1978

(51) INT CL3 C08G 73/02; A61K 7/06, 7/13, 7/40; C08L 79/02; C11D



C3R 35C 35D1 35D2 35D3 35D4 35P2 C11 C12 C16 C25 C29 C4 C6X C9A L2A L2B L2X

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D1B 2A4

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(54) COSMETIC COMPOSITIONS BASED ON **QUATERNISED POLYMERS**

(71) We, L'OREAL, a French Body Corporate of 14, Rue Royle Paris 75008, France, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly

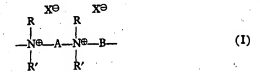
described in and by the following statement:—

The present invention relates to the use in cosmetics of polymers possessing quaternised ammonium groups, to the cosmetic compositions containing these polymers,

and to a process for treating hair or skin using these polymers. Certain cationic polymers wherein the quaternised nitrogen atoms form part of the macro-chain are known and their use as pesticides, as flocculating agents, as surface-active agents and as ion exchangers has already been proposed.

It has now been discovered that, surprisingly, such polymers exhibit valuable

cosmetic properties if they are applied to the hair or to the skin. According to the present invention there is provided a method of treating the hair and/or the skin which comprises applying thereto at least one quaternised polymer comprising recurring units of the general formula I:



in which X9 represents an anion derived from an inorganic or organic acid, R is a lower (C₁ to C₅) alkyl group or a —CH₂—CH₂OH group, R' is an aliphatic radical, an alicyclic radical or an araliphatic radical, which radical contains at most 20 carbon atoms, or two radicals R and R' attached to the same nitrogen atom form, with the latter, a ring which can contain a second hetero-atom other than nitrogen, A represents a divalent group of the formula

$$\begin{array}{c}
-CH_2 - C \\
CH_2 - C \\
CH_2 - CH - (CH_2)_x - CH - (CH_2)_t - CH - (CH_2)_t
\end{array}$$
or $-(CH_2)_x - CH - (CH_2)_x - CH - (CH_2)_t$

in which x, y and t are each independently 0 or an integer which can vary from 1 to

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11 and are such that the sum (x+y+t) is greater than or equal to 0 and less than 18, and E and K represent a hydrogen atom or an aliphatic radical having fewer than 18 carbon atoms, or A represents a divalent group of the formula:

$$\begin{array}{c} -(CH_2)_v - S - (CH_2)_n - \\ -(CH_2)_n - O - (CH_2)_v - \\ -(CH_2)_n - S - (CH_2)_n - \\ -(CH_2)_n - SO - (CH_2)_n - \\ -(CH_2)_n - SO_2 - (CH_2)_n - \\ -(CH_2)_n - SO_2 - (CH_2)_n - \\ \end{array}$$

n being an integer equal to 2 or 3, B represents a divalent group of the formula

or
$$-(CH_2)$$
, $-CH$ — (CH_2) ,

in which D and G represent a hydrogen atom or an aliphatic radical having fewer than 18 carbon atoms, and v, z and u are each independently 0 or an integer which can vary from 1 to 11, and any two of them can simultaneously be 0, but the sum (v+z+u) is greater than or equal to 1 and less than 18 and the sum (v+z+u) is greater than 1 if the sum (x+y+t) is 0, or B represents a divalent group of the formula

20 or $-(CH_2)_n-O-(CH_2)_n$

ve.

n being defined as above.

Certain polymers of the formula I are known whilst others are new (as indicated below).

below).

The polymers of which the recurring units correspond to the formula I will, for simplicity, be referred to by the expression "polymers of formula I".

The terminal groups of the polymers of the formula I vary depending on the relative proportions of the starting reactants. They can be either of the type

N—A—

or-of-the-type

X-B-

In the general formula I, X\(\tilde{\to}\) especially represents a halide, (bromide, iodide or chloride) anion or an anion derived from other inorganic acids, such as phosphoric acid or sulphuric acid, or an anion derived from an organic sulphonic or carboxylic acid, especially an alkanoic acid having 2 to 12 carbon atoms (for example acetic acid), a phenylakanoic acid (for example phenylacetic acid), benzoic acid, lactic acid, citric acid or para-toluenesulphonic acid; the substituent R preferably represents an alkyl group having 1 to 6 carbon atoms; if R' represents an aliphatic radical, it is in particular an alkyl or cycloalkyl-alkyl radical having fewer than 20 carbon atoms and preferably not having more than 16 carbon atoms; if R' represents an alicyclic radical, it is especially a cycloalkyl radical with 5 or 6 ring members; if R, represents an araliphatic radical, it is especially an aralkyl radical, such as a phenylalkyl radical, of which the alkyl group preferably comprises from 1 to 3 carbon atoms; if two radicals R and R' attached to one and the same nitrogen atom form a ring therewith, R and R' can together especially represent a polymethylene radical having 2 to 6 carbon atoms, and the ring can contain a second hetero-atom, for example oxygen or sulphur; if the substituent E, K, D or G is an aliphatic radical, it is especially an alkyl radical having

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1 to 17 carbon atoms and preferably 1 to 12 carbon atoms; v, z and u are preferably from 1 to 5, but two of them can be zero; x, y and t are preferably from 0 to 5; if A or B represents axylylene radical, it can be an o-, m- or p-xylylene radical.

Amongst the polymers of the formula I, the preferred compounds for the cosmetic use according to the invention are especially those for which R is a methyl or hydroxyethyl radical, R' is an alkyl radical having 1 to 16 carbon atoms, a benzyl radical or a cyclohexyl radical, or R and R' together represent the —(CH₂)₅— or

radical; A is a xylylidene radical, a polymethylene radical having 2 to 12 carbon atoms and optionally branched by one or two alkyl substituents having 1 to 12 carbon atoms, a radical of the formula

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or a polymethylene radical having 4 or 6 carbon atoms and containing a hetero-atom grouping of the type of —O—, —S—, —S—S—, —SO— or —SO₂—; B is a xylylidene radical, a polymethylene radical having 3 to 10 carbon atoms and optionally substituted by one or two alkyl radicals having 1 to 12 carbon atoms, a

radical, or a polymethylene radical having 4 or 6 carbon atoms and containing an oxygen hetero-atom; and X is a chlorine, iodine or bromine atom.

It should be noted that the groups A, B, R or R' can have several different values in one and the same polymer I.

Such polymers can be obtained as indicated later in describing the processes of preparation of the polymers of formula I. By way of illustration, the preparation of one such polymer is described in Example 43.

The polymers of formula I can, in particular, be prepared in accordance with the conventional processes reviewed below:

PROCESS 1.

This process consists of carrying out a polyquaternisation reaction of a di-tertiary diamine of the formula

with a dihalide of the formula $X-B_1-X$, in which R, R' and X have the meanings indicated above; A_1 represents A if B_1 represents B, and A_1 represents B if B_1 represents A, the groups A and B being as defined above.

This process for the preparation of the polymers of the formula II can thus be carried out in accordance with one of the two following methods:

PROCESS 1A.

PROCESS 1b.

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If A represents one of the following groups:

$$\begin{array}{cccc} & -(CH_2)_n - S - (CH_2)_n -, \\ & -(CH_2)_n - SO - (CH_2)_n -, \\ & -(CH_2)_n - SO_2 - (CH_2)_n -, \\ & -(CH_3)_n - S - S - (CH_2)_n -, \\ & -(CH_3)_n - S - (CH_2)_n -, \\ & -(CH_3)_n - (CH_3)_n -, \\ & -(CH_3)_n -, \\ &$$

Process 1a is preferably used.

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The polyquaternisation reaction is carried out, for example, in a solvent or a mixture of solvents which favours quaternisation reactions, such as water, dimethylformamide, acetonitrile and the lower alcohols, especially the lower alkanols such as methanol.

The reaction temperature is suitably from 10° to 150° C. and preferably from

to 100° C. The reaction time depends on the nature of the solvent, on the starting reactants

and on the desired degree of polymerisation. In general, the starting reactants are reacted in equimolecular amounts but it is also possible to use either the diamine or the dihalide in a slight excess, this excess being generally less than 20 mol%.

The resulting polycondensate can be isolated at the end of the reaction, either by

filtration or by concentrating the reaction mixture.

It is possible to regulate the average chain length by adding, at the start or in the course of the reaction, a small amount (e.g. 1 to 15 mol% relative to one of the reactants) of a monofunctional reactant such as a tertiary amine or a monohalide. In this case, at least a part of the terminal groups of the resulting polymer I consists either of the tertiary amine group used or of the hydrocarbon group of the monohalide. Examples of limiting the chain length by adding varying amount of triethylamine are given later (see Examples 47 to 49).

This invention includes the use of the polymers of the formula I which have such

In place of the starting reactant it is also possible to use either a mixture of diterminal groups. tertiary diamines or a mixture of dihalides or a mixture of di-tertiary amines and a mixture of dihalides, provided the ratio of the total molar amounts of diamines and dihalides is about 1. This possibility is illustrated later in Example 43.

PROCESS 2. This process consists of subjecting a w-halogenated tertiary amine of the formula

to a self-polyquaternisation reaction.

In this case, the polymer obtained corresponds to the general formula I, in which B=A, with R, R' and X having the meanings mentioned above, and A can only assume the meaning

the sum (x+y+t) having to be greater than 4.

The reaction can be carried out either without solvent or with the same solvents as in Process I, using the same ranges of reaction temperature. As before, it is possible to add a monofunctional reagent to regulate the degree of polyquaternisation. It is also possible to use a mixture of several tertiary w-halogenated amines.

In the two processes for the preparation of the polymers of the formula I mentioned above, the polymer which results can be isolated at the end of the reaction either by filtration or by concentrating the reaction mixture and crystallising the product, if appropriate, by adding a suitable anhydrous organic liquid, for example acetone.

The di-tertiary diamines used as starting products in the above Process 1 can be prepared in accordance with one of the methods indicated below.

Method 1

A primary amine of the formula R—NH₂ is reacted with a dihalide of the formula Hal—A₁—Hal, Hal being a halogen atom and preferably a bromine or iodine atom. The process is suitably carried out at 50° to 150° C., using an excess of primary amine, generally 2 to 5 mols of primary amine per mol of dihalide. After adding to the reaction mixture a basic solution, for example a solution of sodium hydroxide or potassium hydroxide, the di-secondary diamine of the formula

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R—NH—A₁—NH—R

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is obtained. The latter is then subjected to an alkylation reaction according to known methods, the expression "alkylation" here denoting the replacement of a hydrogen atom linked to the nitrogen by a group R' as defined above. The di-tertiary diamine of the formula

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can then be isolated in accordance with the usual methods.

Method 2: (applicable to the case wherein A₁ is a xylyhidene radical or an alkylene radical).

A primary amine R.—NH₂ is reacted with an arylsulphonyl halide Ar—SO₂—Hal, Ar being an aryl group, for example a phenyl or tolyl group, and Hal being a halogen atom, for example a chlorine atom. A sulphonamide of formula Ar—SO₂—NHR is obtained, which is subjected to an alkylation reaction in accordance with known methods for producing a sulphonamide of the formula

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which, on acid hydrolysis, for example by means of an aqueous solution of sulphuric acid, gives the secondary amine of the formula

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The latter is then reacted with a dihalide of the formula Hal—A₁—Hal (Hal and A₁ being defined as above), in the presence of a tertiary amine such as N-ethyldiiso-propylamine, using at least two mols of secondary amine and of tertiary amine per mol of dihalide. The reaction is preferably carried out without a solvent and at a temperature of 50° to 130° C. At the end of the reaction, the mixture is taken up in water to dissolve the amine salts and the solution is extracted with a suitable solvent, for example ethyl acetate. The extracts are then washed with an aqueous alkaline solution (sodium hydroxide or potassium hydroxide), and then with water. The organic phase is then dried, after which the di-tertiary diamine is isolated either by distillation or by concentration under reduced pressure.

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Method 3:

A primary amine RNH₂ is reacted with an arylsulphonyl halide, as indicated in Method 2. The sulphonamide obtained, of the formula Ar—SO₂—NHR, is reacted with a dihalide of the formula Hal—A₁—Hal at a temperature of, say, 80° to 140° C.

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The disulphonamide formed, of the formula

can be subjected directly, without being isolated, to acid hydrolysis. This acid hydrolysis can be carried out, for example, in an 85% strength aqueous solution of sulphuric acid, at a temperature of, say, 120 to 145° C., for a period of, say, 7 to 20 hours. The di-secondary diamine of the formula

is obtained, and is subjected to an alkylation reaction, in accordance with known methods, so as to form the di-tertiary diamine of the formula

This method 3 is particularly useful in the case wherein $A_1 = (CH_2)_5$, because it avoids the cyclisation side-reactions observed if methods 1 and 2 are used.

Method 4:
A secondary amine of the formula

H—N R

is reacted with a dihalide of the formula Hal—A₁—Hal (Hal and A₁ being defined as indicated above) in the presence of an alkali metal carbonate or a tertiary amine such as N-ethyldiisopropylamine.

If the reaction is carried out in the presence of an alkali metal carbonate, at least one mol of carbonate is used per mol of dihalide and ethanol is generally used as the solvent; at the end of the reaction, the inorganic salts are filtered off, the ethanol is driven off under reduced pressure and the reaction products are separated by distillation

When the reaction is carried out in the presence of a tertiary amine, at least two mols of secondary amine and tertiary amine are used per mol of dihalide; the reaction is preferably carried out without a solvent and at a temperature of 50° to 130° C.; at the end of the reaction, the mixture is taken up in water to dissolve the amine salts and the solution is extracted with a suitable solvent, for example ethyl acetate; the extracts are then washed with an aqueous alkaline solution (sodium hydroxide or potassium hydroxide) and then with water; the organic phase is then dried and thereafter the di-tertiary diamine is isolated, for example by distillation or by concentration under reduced pressure.

The secondary amine starting materials (with R'= -CH₂-CH₂-OH) can be prepared by reaction of a halogen derivative R-Hal with monoethanolamine.

The secondary amine starting materials (wherein R' is not —CH₂—CH₂—OH) are obtained, for example, by reacting a primary amine of the formula R—NH₂ with an aryisulphonyl halide Ar—SO₂—Hal, Ar being an aryl group, for example a phenyl or tolyl group, and Hal being a halogen atom, for example a chlorine atom. A sulphonamide of the formula Ar—SO₂—NHR is obtained, which is subjected to an alkylation reaction in accordance with the known methods, so as to form a sulphonamide of the formula

which, on acid hydrolysis, for example with an aqueous solution of sulphuric acid, gives the secondary amine of the formula

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5 Method 5: (applicable to the case wherein A₁ represents —CH₂—CHOH—CH₂—)

Epichlorohydrin is reacted with a secondary amine R—NH—R', preferably used in excess. The reaction is suitably carried out in a solution or suspension in water at a temperature of 40° to 100° C., with 3 to 10 mols of secondary amine per mol of epichlorohydrin. After adding to the reaction mixture a basic solution, for example a solution of sodium hydroxide or potassium hydroxide, and extracting with an appropriate solvent (for example ethyl acetate), a mixture of secondary amine starting material and di-tertiary diamine is obtained, which is separated by distillation.

Method 6: (applicable to the case wherein A1 represents

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$$-(CH_2)_a$$
 $-S$ $-S$ $-(CH_2)_a$ $-)$

An alkali metal thiosulphate is reacted with an amine of the formula

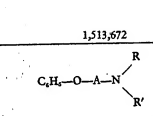
to form a Bünte salt, which is hydrolysed to the disulphide by means of a solution of sodium hydroxide or potassium hydroxide. The reaction is suitably carried out in water at a temperature of 40° to 100° C. When the Bünte salt has been formed completely, it is hydrolysed and the corresponding disulphide is extracted in an appropriate solvent such as ethyl acetate. The solvent can be removed by distillation under reduced pressure and the di-tertiarydiamine isolated and purified, if desired, by distillation under reduced pressure.

The w-halogenated tertiary amines used as starting materials in Process 2 described above can themselves be prepared by applying processes described by M. R. LEHMAN, C. D. THOMPSON and C. S. MARVEL, J.A.C.S., 55, 1977 (1933) and by LITTMANN and MARVEL, J.A.C.S., 52, 287 (1930), replacing the dimethylamine starting material, as required, by the appropriate secondary amine of the formula



and R' being defined as above. They can also be obtained by the reaction, in the presence of a proton acceptor such as N-ethyldiisopropylamine, of a compound of the formula Br—A—OC, H,, with a secondary amine of the formula

in which A, R and R' are defined as above, followed by reaction with hydrobromic acid of the resulting compound of the formula



	, R	•
5	under the conditions described in the two J.A.C.S. references referred to above. Though the invention is not limited to the use of polymers I having a degree of polymerisation varying within a particular range, it can be said that the polymers of the formula I which are most generally suitable have a molecular weight which is	. 5
	from 5,000 to 50,000. They are generally soluble in at least one of the three following solvents: water, ethanol or a water-ethanol mixture. It is possible to obtain films, which in particular exhibit good affinity to hair, by	10
10	evaporation of the solution of the polymer. As indicated above, the polymers of the formula I have valuable cosmetic properties which permit their use in the preparation of cosmetic compositions. Such cosmetic compositions comprise polymers of the formula I either by way of	
15	the main active ingredient or by way of an additive. These cosmetic compositions can be in the form of, for example, aqueous, alcoholic or aqueous-alcoholic solutions, (the alcohol being, in particular, a lower alkanol such as ethanol or isopropanol) or in the form of creams, gels or emulsions, or in the form of aerosol cans also containing a propellant. Adjuvants generally present in the cosmetic compositions of the invention are,	15
20	for example, perfumes, dyestuffs, preservatives, sequestering agents and thickeners. The compositions suitable for application to the hair and/or skin which comprise at least one cosmetic adjuvant which is: (i) a cosmetic resin such that the composition is in the form of a hair lacquer or wave-setting lotion;	20
- 25	 (ii) a hair dye; (iii) a hair restructuring agent; (iv) a surface-active agent such that the composition is in the form of a shampoo; or (v) an oil such that the composition is in the form of an emulsion for application 	30
30	It should be noted that the cosmetic compositions according to the invention can be either ready-to-use compositions or concentrates which have to be diluted before use. Hence, the cosmetic compositions of the invention are not restricted to a posterior of the cosmetic composition of the formula I.	•
35	In general, in the cosmetic compositions of the invention, the concentration of polymers of the formula I is from 0.5 to 10% by weight, preferably from 0.5 to 5% by weight. The polymers of the formula I in particular have valuable cosmetic properties	35
40	Thus, if they are applied to the head of hair either by themselves of together with other active substances, in the course of a treatment such as shampooing, dyeing or provesting they noticeably improve the properties of the hair.	40
45	For example, they assist the treatment and facilitate the combing out of wer harr. Even at a high concentration, they do not make wet hair sticky to the touch. In contrast to the customary cationic agents, they do not make dry hair heavy and consequently facilitate bouffant hair styles. They make dry hair springy and give it a	45
50	They contribute efficiently to removing the defects of hair which has been sensitised by treatments such as bleaching, permanent waving or dyeing. It is known, in fact, that sensitised hair is frequently dry, dull and rough, and difficult to comb out and set. They are in particular of great value when they are used as pretreatment agents,	50
55	which is itself followed by an anionic and/or non-ionic shampoo or before an oxidation dyeing which is itself followed by an anionic and/or non-ionic shampoo. The hair is then particularly easy to comb out and feels very soft. They can also be used as pretreatment agents in other operations involving the resument of hair for example permanent wayings.	55
60	The cosmetic compositions for application to the hair can be in the form of, for example, aqueous, alcoholic or aqueous-alcoholic solutions (the alcohol generally being a lower alkanol such as ethanol or isopropanol), or in the form of creams, gels	60

	or emulsions, or in the form of sprays. They can also be packaged in aerosol containers, which also contain a propellant, for example nitrogen, nitrous oxide or chloro-	•
	fluorinated hydrocarbon of the "Freon" [Registered Trade Mark] type. The adjuvants generally present in the cosmetic compositions for hair include	
5	perfumes, dyestuffs, preservatives, sequestering agents, thickeners and emulsifiers or resins usually employed in cosmetic compositions for hair (cosmetic resins). The polymers of the formula I can be present in, for example, wavesetting lotions,	5
	treatment lotions and setting creams or gels, or as an additive in shampoo composi- tions, wavesetting compositions, permanent waving compositions, dyeing compositions,	
10	restructuring lotions, treatment lotions for counteracting dandruff, or hair lacquers. The cosmetic compositions for hair according to the invention or used in the method of this invention, are thus in particular:	10
	a) treatment compositions which comprise, by way of active ingredient, at least	
45	one polymer of the formula I in aqueous or aqueous-alcoholic solution.	15
15	The content of polymer of the formula I is generally from 0.5 to 10% by weight and preferably from 0.5 and 5% by weight.	7.2
	The pH of these solutions is approximately 7 and can vary, for example, from	
	6 to 8. It is possible, if necessary, to bring the pH to the desired value by adding either an acid such as citric acid or a base, especially an alkanolamine such as mono-	
20	ethanolamine or triethanolamine.	20
	In order to treat hair with such a lotion, the lotion is applied to wet hair and is	
	allowed to act for, say, 3 to 15 minutes, and then the hair is rinsed. If desired, a conventional wavesetting operation can then be carried out;	
	b) shampoos which comprise at least one polymer of the formula I and a cationic,	
25	non-ionic or anionic detergent. The cationic detergents are, especially, long-chain quaternary ammonium com-	25
	pounds, esters of fatty acids and amino-alcohols, or polyether-amines.	
	The non-ionic detergents are especially esters of polyols and sugars, products	. •
30	resulting from the condensation of ethylene oxide with fatty compounds, with long chain alkyl-phenols, with long chain mercaptans or with long chain amides, and	30
50	polyhydroxylic polyethers of a fatty alcohol.	
	The anionic detergents are especially alkali metal salts, ammonium salts or amine or amino-alcohol salts of fatty acids such as oleic acid, ricinoleic acid and acids from	
	copra oil or from hydrogenated copra oil; alkali metal salts, ammonium salts or	
35	amino-alcohol salts of fatty alcohol sulphates, and especially of C ₁₅ —C ₁₆ and C ₁₆ fatty alcohols; alkali metal salts, magnesium salts, ammonium salts or amino-alcohol salts of oxyethyleneated fatty alcohol sulphates; products resulting from the condensa-	35
	tion of fatty acids with isethionates, with taurine, with methyltaurine or with sarcosine;	
40	alkylbenzene-sulphonates, especially with a C ₁₂ alkyl group; and alkylarylpolyether-	40
40	sulphates or monoglyceride-sulphates. All these anionic detergents, as well as numerous others not mentioned here, are well known and are described in the literature.	40
•	These shampoos can also contain various adjuvants, for example perfumes, dye-	
	stuffs, preservatives, thickeners, foam stabilisers, softening agents or one or more cosmetic resins.	
45	In these shampoos, the concentration of detergent is generally from 5 to 50%	45
	by weight and the concentration of polymer of the formula I or I ₄ is from 0.5 to 10%, and preferably from 0.5 to 5% by weight.	
	c) wavesetting lotions, especially for sensitised hair, which comprise at least one	
60	polymer of the formula I, in aqueous, alcoholic or aqueous-alcoholic solution.	50
50	They can also contain another cosmetic resin. The cosmetic resins which can be used in such lotions are very varied and are especially vinyl or crotonyl homopoly-	50
	mers or copolymers, for example polyvinylpyrrolidone, copolymers of vinylpyrrolidone	
	and vinyl acetate and, copolymers of crotonic acid and vinyl acetate. The concentration of the polymers of the formula I in these wavesetting lotions	
55	generally varies from 0.5 to 5% and preferably from 0.5 to 3% and the concentration	55
	of the other cosmetic resin varies essentially within the same proportions.	
	The pH of these wavesetting lotions is generally from 3 to 9 and preferably from 4.5 to 7.5. If desired, the pH can be changed, for example by adding an alkanol-	
	amine such as monoethanolamine or triethanolamine;	
60	d) dyeing compositions for hair, which comprise at least one polymer of the formula I, a dyeing agent and a carrier.	60
	The carrier is preferably chosen so as to form a cream.	
	The concentration of the polymers of the formula I in these dyeing compositions	
65	is generally from 0.5 to 15% by weight and preferably from 0.5 to 10% by weight. In the case of oxidation dyeing, the dyeing composition can be packaged in two	65
	,	

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••	1,513,672	10
10	parts, the second part comprising hydrogen peroxide. The two parts are mixed at the	
	time of use. Examples of such compositions and of their use are given in the working	
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5	e) hair lacquers, which comprise an alcoholic or aqueous-alcoholic solution of a customary cosmetic resin for lacquers, and at least one polymer of the formula I, customary cosmetic resin for lacquers, and at least one polymer of the formula I,	. •
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	tion by adding the customary cosmetic resin and the polymer of the formula I to a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an anhydrous aliphatic alcohol such as ethanol or isopropanol and a mixture of an annotation and a mixture of an analysis aliphatic alcohol such as ethanol or isopropanol and a mixture of an analysis aliphatic alcohol and a mixture of an analysis aliphatic alcohol and a mixture of an analysis	10
10	1:	
	generally from 0.5 to 3% by weight and the concentration of the polymer of the formula I is generally from 0.5 and 3% by weight.	15
15	Of course, it is possible to add adjuvants such as dyestuffs, plasticisers or any other	
	f) restructuring treatment lotions, which comprise at least one agent possessing hair restructuring properties and at least one polymer of the formula I.	'
20	THE PARTY OF ACCOUNT WHICH CAN BE HISPET IN SHELL TOURISH AND	20
20	methylol derivatives described in French Patents No. 1,515,575, 2,525,503, 25	
	7,519,982 and 1,527,085.	
	to 10% by weight and the concentration of the polymer of the formula I is generally	25
25		23
	g) Pretreatment compositions especially in the form of aqueous or aqueous- alcoholic solutions, optionally in an aerosol container, or in the form of creams or gels,	
	the second compositions being intended to be applied to the sum	
,		30
30	oxidation dyeing followed by an anionic and/or non-ionic shampoo, or before a permanent waving treatment.	
	T	
25	5% by weight. The pH of these compositions which is near 7, generally from 3 to 9 and especially from 6 to 8.	35
35	There mentment compositions can contain various aujuvaits (101 onthips	
	resins) usually employed in cosmetic compositions for hair including agents for changing the pH (for example, amino-alcohols such as monoethanolamine) as indicated	
		40
40	The polymers of the formula I also possess valuable cosmette properties	. 40
,,,	they are applied to the skin. In particular, they assist moisturising of the skin and thus prevent it from drying	
	The second of the chiral considerable southess to the touch	
	The accomption compositions for the skin can be presented, for example in the	45
45	form of creams, gels, emulsions or aqueous, alcoholic or aqueous-alcoholic solutions. The concentration of the polymer of the formula I in these compositions for the	
	1 to the managed list factor (1 S to 1007) by Welton.	
	The editionts generally present in these cosmetic compositions are, for example,	
	perfumes, dyestuffs, preservatives, thickeners, sequestering agents and emulsifiers. These compositions for the skin are, in particular, treatment creams or lotions for	50
50	the hands or face, anti-sunburn creams, tinted creams, cleaning minks, and rouning	
	Last liquide on (are present) in decourant compositions.	•
	These compositions can be prepared in accordance with the usual methods. For example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example, in order to prepare a cream, it is possible to emulsify an aqueous for example and the contract of the	
55	phase containing a solution of the polymer I and optionally other ingredients or	55
33	11 and an aily where	
	The oily phase can consist of various products such as liquid paraffin, "Vaseline" [Registered Trade Mark] oil, sweet almond oil, avocado oil, olive oil	
	and a state of factor acide cuch as divicety manasterials, sully of footbody partitions	60
60	and alkyl myristates such as propyl, butyl of cetyl myristate. It is also possible to any	00
	fatty alcohols such as cetyl alcohol or waxes such as beeswax. The polymers of the formula I can be present in the cosmetic compositions for	
•	at a line of they as an additive of as the main active illerement in treatment creams	
	or lotions for the hands or face, or as an additive in compositions of and summer	65
65	creams, tinted creams, cleansing milks and foam bath liquids.	

	The preferred polymers of the formula I are those described below in Examples	
5	No. 1 to 140. In a particular embodiment of the invention the method for the treatment of hair consists of applying to the hair, before an anionic shampoo, or before an oxidation dyeing treatment followed by an anionic shampoo, at least one polymer of the formula I by means of a pretreatment composition as defined above. The following Examples further illustrate the present invention.	5
	EXAMPLES OF THE PREPARATION OF THE STARTING DIAMINES. Preparation No. 1.	4.
10	1,6-N,N'-Dibutyl-N,N'-dimethyl-diamino-hexane 61 g of 1-dibromohexane is added slowly, whilst stirring, to 91 g of n-butylamine previously brought to a temperature of 75° C. The temperature of the reaction mixture actually rises to 95° C. When the introduction is complete, the hydrobromide formed	10
15	crystallises and the temperature rises to 110° C. The mixture is cooled to 60° C. and 250 cm³ of water and 50 cm³ of a concentrated aqueous solution of sodium hydroxide are introduced successively. Stirring is continued for ½ hour and then the precipitate of 1.6-N,N'-dibutyl-diamino-hexane is filtered off and dried under reduced pressure.	15
20	b) 37 g of the latter product are added, with stirring and whilst keeping the temperature below +5° C, to a mixture of 108 g of pure formic acid and 11 g of water. 117 g of a 30% strength aqueous solution of formaldehyde are then introduced over the course of 10 minutes. The temperature is then raised slowly to 100° C. and	20
25	the mixture is stirred at this temperature until carbon dioxide ceases to be evolved. The reaction mixture is then concentrated under reduced pressure and the residue is rendered alkaline by adding approximately 150 cm³ of a concentrated aqueous solution of sodium hydroxide. The mixture is extracted three times with 200 cm³ of isopropyl ether. The combined organic phases are washed three times with 100 cm³ of water, dried and concentrated. The residue is distilled in vacuo and 72 g of 1,6-N,N'-dibutyl-N,N'-dimethyl-diamino-hexane which distils at 98—99° C. under 0.1 mm of mercury, are collected.	25
30	Preparation No. 2. 1,3-N,N'-Dimethyl-N,N'-dioctyl-diamino-propane.	30
35	a) N-Methyl-octylamine. 523 g of octylamine benzenesulphonamide are dissolved in 1,500 cm ² of anhydrous xylene. 835 cm ³ of a 2.4 N solution of sodium ethylate in ethanol are then introduced, with stirring, and the ethanol is then removed by distillation. Whilst keeping the stirred reaction mixture at 100—110° C., 385 cm ³ of methyl sulphate are introduced over the course of 1 hour and the mixture is heated under reflux for 4 hours. After cooling,	35
40	the inorganic salts are filtered off. 1,500 cm ³ of a concentrated aqueous solution of sodium hydroxide are added to the filtrate, decanting is effected and the xylene phase is washed 4 times with 1,000 cm ³ of water and then concentrated. The residue obtained is added to a mixture of 1,400 g of concentrated sulphuric	40
45	acid and 560 g of crushed ice and the whole is heated, with stirring, to 160° C. for 16 hours. After cooling, the reaction mixture is poured onto 3 kg of crushed ice and rendered alkaline by adding 3,500 cm³ of a concentrated aqueous solution of sodium hydroxide. The mixture is extracted three times with 2,000 cm³ of ethyl acetate, and the organic phases are washed with water, dried and concentrated under reduced pressure. The residue is distilled and the fraction which distils at 45—50° C. under 0.2 mm of mercury is collected.	45
50	b) 1,3-N,N'-Dimethyl-N,N'-dioctyl-diamino-propane. 69 g of 1,3-dibromo-propane are added to a mixture of 107 g of N-methyl- octylamine and 87.5 g of N-ethyl-diisopropylamine whilst keeping the temperature at 100—105° C. The mixture is then stirred for 7 hours at 120° C. and is cooled, and 500	50
55	cm ³ of water and 200 cm ³ of ether are added. The aqueous phase is decanted and the ether phase is treated with 50 cm ³ of a concentrated aqueous solution of sodium hydroxide, decanted and washed three times with 100 cm ³ of water. After drying, the ether is evaporated and then the starting N-methyl-octylamine which has not reacted. The residue obtained comprises two phases which are separated.	55
60	The clear upper phase is purified by distillation; 1,3-N,N'-dimethyl-N,N'-dioctyl-diamino-propane, which distils at 150—153° C. under 0.5 mm of mercury, is collected.	60
	.•	

12		
'	Preparation No 3. 1,5-N,N'-Didecyl-N,N'-dimethyl-diamino-pentane.	
5 .	a) 1,5-N,N-Didecyl-diamino-pentane. 297 g of n-decylamine benzenesulphonamide are reacted with sodium ethylate in 297 manner analogous to that described in Example 2, in order to form the correspondant manner analogous to that described in Example 2, in order to form the correspondant manner analogous to that described in Example 2, in order to form the correspondant in the correspon	5
40	diiodopentane are introduced, with shiring the xylene for 4 hours and is then cooled, mixture is kept at the reflux temperature of the xylene for 4 hours and is then cooled, and 500 cm ³ of water are added with stirring in order to dissolve the inorganic salts	10
15	formed. The organic phase is then decanted, dried and concentrated under reduced pressure. The residue obtained is heated for 14 hours at 130° C. in the presence of 300 cm³ of 85% strength sulphuric acid. After cooling, the reaction mixture is poured onto 1.5 kg of crushed ice and is brought to pH 10 by adding a 30% strength aqueous sodium hydroxide solution. The precipitate formed is filtered off, washed with water and dried. 1,5-N,N'-Didecyl-diamino-pentane is obtained.	15
20	b) 1,5-N,N'-Didecyl-N,N'-dimethyl-diamino-pentane. Methylation of the product obtained under a), in accordance with a method analogous to that described in Preparation No. 1, gives 1,5-N,N'-didecyl-N,N'-dimethyl-diamino-pentane. Boiling point=193—195° C. (0.4 mm Hg).	20
25	Preparation No. 4. 1,3-N,N'-Didodecyl-N,N'-dimethyl-diamino-propan-2-ol. 26.6 g of epichlorohydrin are added slowly to a vigorously stirred mixture of 370 g of N-methyl-dodecylamine and 600 cm ³ of water. At the end of the addition, the reaction mixture is heated to 90° C. for 12 hours. After cooling, 10 cm ³ of a concentrated aqueous sodium hydroxide solution are added and the mixture is extracted three times with 200 cm ³ of ethyl acetate. The extraction solutions are dried over sodium times with 200 cm ³ of ethyl acetate. The extraction solutions are dried over sodium times with 200 cm ³ of ethyl acetate. The extraction solutions are dried over sodium	25
30	sulphate and concentrated under reduced pressure. A first fraction corresponding to the excess N-methyl-dodecylamine is obtained; the second fraction, distilling at 235°—240° C. under 1.5 mm Hg is the di-tertiary diamine.	30
35	Preparation No. 5. 1,3-N,N'-Di-2-hydroxyethyl-N,N'-dioctyl-diamino-propane. A mixture consisting of 100 g of N-2-hydroxyethyloctylamine, 28.6 g of 1,3-dibromopropane, 22 g of potassium carbonate and 300 cm ³ of ethanol is heated under reflux for 50 hours. The inorganic salts are removed by filtration, the ethanol is distilled under reduced pressure and the residue is then distilled in vacuo. The expected diamine distills at 200—206° C. under 1 mm of mercury.	35
40	Preparation No. 6. N-Butyl-N-methyl-2-aminoethyl disulphide. 44 g of N-butyl-N-methyl-2-bromoethylamine hydrobromide are dissolved in 10 cm³ of water. Thereafter the pH of the solution is adjusted to 7 by adding dilute	40
45	sodium hydroxide whilst keeping the temperature at about of C. In temperature is then raised to 60° C. whilst introducing a solution of 43.7 g of sodium thiosulphate pentahydrate in 15.6 cm³ of water. The reaction mixture is kept at 60° C. for 8 hours and then cooled, after which	45
50	is left to stand for 2 hours, the organic phase is then extracted with for the organic phase is then extract a with for the organic phase is then extract a with for the organic phase is then extracted with for the organic phase is then extract in the organic phase is then extract phase is the organic phase in the organic phase is the extract phase is the organic phase is the extract phase is the organic phase is the organic phase is the extract phase is the organic phase is the extract phase is the organic phase is the organic phase is the extract phase is the organic phase is the extract phase is the organic phase is the organic phase is the organic phase is the organic phase is the extract phase is the organic phas	50
55	The di-tertiary diamines indicated in Table I which follows were prepared analogously in accordance with the methods described previously; these di-tertiary diamines are used as starting materials in the examples of the preparation of polymers of the formula I, given later in the experimental part.	55

TABLE I

		TABLE I		
Preparation No.	Method No.	A,	R	R'
7	2	(CH ₂) ₁₀	СН,	C,,H,,
8	1	(CH ₂),	CH,	C ₄ H ₉
9	2	(CH ₂) ₆	CH,	C,H,7
10	· 2	(CH ₂) ₁₀	СН,	C,H,,
11	1	(CH ₂) ₁₀	CH,	C ₄ H ₉
12	2	(CH ₂) ₆	Сн,	C, H,,
. 13	1	(CH ₂) ₆	CH ₃	cyclohexyl
14	1	(CH ₂),	CH,	isobutyl
15	2	(CH ₂),	CH,	C,,H,,
. 16	2	(CH ₄) ₆	CH,	C,2H,5
17	, 1	p-xylylene	CH,	C.H.
18	1	m-xylylene	CH,	C₄H,
19	1	(CH ₂) ₁₂	СН,	C.H.
20	• 1	(CH ₂) ₆	CH,	С,Н,
21	1	(CH ₂) ₆	CH,	isopropyl
22	1	(CH ₂),	СН,	benzyl
23	1	_(CH,)_CH_	СН,	C,H,,
		CH,		
24	· 1	(CH ₂) ₈	CH,	isobutyl
		С.Н. С.Н.		
25	1	HC_(CH ₂) _s _CH	СН,	C,H,,
		Сн, Сн,		
26	1	_СН,_СН_СН,_	СН,	C ₄ H ₉
		C ₁₂ H ₂₈		.]
27	٠.	сн,_сн_сн,	СН,	C,0H21
•	, ,	ОН		
28		Сн, Сн_Сн,	CH,	C _s H ₁ ,
		ОН		
		·	: . I	

TABLE I (continuation)

Preparation No.	Method No.	$\mathbf{A_i}$	R	R'
29	*	(CH ₂),	_CH,CH,OH	C,2H25
30		(CH ₂) ₆	_CH,CH,OH	C,2H25
31		$(CH_2)_6$	_CH,CH,OH	C ₄ H ₉
32		(CH ₂) ₂ _S_S_(CH ₂),	СН,	C,2H25
33		(CH ₂) ₂ -S-S-(CH ₂) ₂	CH ₃	C ₆ H ₁₇

a vacuum (of the order of 0.1 mm of mercury) in the presence of phosphorus 5 pentoxide. In Examples 1 to 73, the process used is Process 1a. Polymer of the formula I (with $R=R'=CH_3$, $A=(CH_2)_{\omega}$ $B=(CH_2)_3$ and X=Br) A solution consisting of 172.3 g of N,N,N',N'-tetramethyl-hexamethylenediamine and 202 g of 1,3-dibromopropane in 650 cm³ of a 50:50 mixture of methanol and 10 10 dimethylformamide is stirred for 170 hours at ambient temperature. A white precipitate is obtained on adding anhydrous acetone; it is filtered off dried. The polymer obtained contains 35.4% of Br. EXAMPLE 2. Polymer of the formula I (with R=R'=CH₈, A=B=(CH₂), and 15 15 X=Br)A solution of 172.3 g of N,N,N',N'-tetramethyl-hexamethylenediamine and 244 g of 1,6-dibromohexane in 1,600 cm3 of anhydrous methanol is heated under reflux for 24 hours, whilst stirring. 20 The polymer obtained contains 36.6% of Br. 20 It is water-soluble. EXAMPLE 3. Polymer of the formula I (with R=R'=CH₂, A=(CH₂)₂, B= and X=Br) A mixture of 116.2 g of N,N,N',N'-tetramethyl-ethylenediamine and 264 g of pxylylidene bromide in 3,200 cm³ of anhydrous methanol is heated under reflux for 50 25 hours. On-cooling, a precipitate is obtained, which is filtered off and dried. The resulting polymer contains 38.8% of Br. It is soluble in water-ethanol mixtures. EXAMPLE 4. Polymer of the formula I (with R=CH₃, R'=C₁₂H₂₅, A=(CH₂)₃, B=(CH₂)₄ and X=Br) 30 30 A solution of 438 g of N,N'-didodecyl-N,N'-dimethyl-trimethylenediamine and 244 g of 1,6-dibromohexane in 3,200 cm³ of anhydrous methanol is heated under reflux for 80 hours. 35 35 The resulting polymer contains 23.4% of Br. It is soluble in ethanol. Polymer of the formula I (with $R=CH_3$, $R'=C_4H_9$, $A=(CH_2)_6$, EXAMPLE 5. B=(CH₂)₁₀ and X=Br).

A solution of 256 g of N,N'-dibutyl-N,N'-dimethyl-hexamethylenediamine and 40 300 g of 1,10-dibromodecane in 3,200 cm³ of anhydrous methanol is heated under

The polymer obtained contains 25.0% of Br. It is soluble in water and in ethanol.

5	EXAMPLE 6. Polymer of the formula I (with R=CH ₃ , R'=C ₈ H ₁₇ , A=(CH ₂) ₁₀ , B=(CH ₂) ₄ and X=Br) A solution consisting of 424 g of N,N'-dimethyl-N,N'-dioctyl-decamethylenediamine and 216 g of 1,4-dibromobutane in 3,200 cm ³ of anhydrous methanol is heated under reflux for 60 hours. The resulting polymer contains 21.1% of Br.	5
10	EXAMPLE 7. Polymer of the formula I (with R=CH ₃ , R'=C ₈ H ₁ ., A=(CH ₂) ₅ , B=(CH ₂) ₄ and X=Br) A solution consisting of 326 g of N,N'-dimethyl-N,N'-dioctyl-trimethylenediamine and 216 g of 1,4-dibromobutane is heated under reflux for 60 hours. The polymer obtained contains 26.0% of Br.	10
15	EXAMPLE 8. Polymer of the formula I (with R=CH ₃ ,R'=C ₁₂ H ₂₃ , A=(CH ₂) ₁₀ , B=(CH ₂) ₄ and X=Br) A solution consisting of 537 g of N,N'-didodecyl-N,N'-dimethyl-decamethylene-diamine and 216 g of 1,4-dibromobutane in 3,200 cm ⁶ of anhydrous methanol is heated under reflux for 80 hours. The polymer obtained contains 20.6% of Br. It is soluble in ethanol.	15
20	EXAMPLE 9. Polymer of the formula I (with R=R'=CH ₈ , A=B=(CH ₂) ₁₀ and X=Br) A solution obtained from 225 g of N,N,N',N'-tetramethyl-decamethylenediamine and 301 g of 1,10-dibromodecane in 3,200 cm ⁶ of anhydrous methanol is heated under reflux for 25 hours. The polymer obtained contains 28.0% of Br.	20
25	EXAMPLE 10. Polymer of the formula I (with R=CH ₃ , R'=C ₉ H ₁ ,A=B=(CH ₂). X=Br) A solution consisting of 368 g of N,N'-dimethyl-N,N'-dioctyl-hexamethylenediamine and 244 g of 1,6-dibromohexanc in 3,200 cm ³ of anhydrous methanol is heated	. 25
30	under reflux for 40 hours. The polymer obtained contains 24.3% of Br. EXAMPLE 11. Polymer of the formula I (with R=CH ₃ , R'=C ₄ H ₅ , A=(CH ₂) ₃ , B= CH ₂ CH ₂ and X=Br)	30
35	A mixture of 214 g of N,N'-dibutyl-N,N'-dimethyl-trimeta, g of p-xylylidene bromide in 1,800 cm ³ of methanol is heated under reflux for 65 hours. The polymer obtained contains 30.7% of Br. It is soluble in water and in water-ethanol mixtures.	35
40	EXAMPLE 12. Polymer of the formula I (with R=CH ₃ , R'=C ₄ H ₉ , A=B=(CH ₂) ₁₀ and X=Br) A solution of 312 g of N,N'-dibutyl-N,N'-dimethyl-decamethylenediamine and 301 g of 1,10-dibromodecane in 3,200 cm ³ of anhydrous methanol is heated under reflux for 55 hours. The polymer obtained contains 23.2% of Br. It is soluble in water and in ethanol.	40
45	EXAMPLE 13. Polymer of the formula I (with R=R'=CH ₂ , A=(CH ₂) ₄ , B=(CH ₂) ₄ and X=Br) A solution consisting of 172.3 g of N,N,N',N'-tetramethyl-hexamethylenediamine and 216 g of 1,4-dibromobutane in 650 cm ³ of a 50:50 mixture of methanol and	45
50	dimethylformamide is stirred for 170 hours at ambient temperature. On addition of anhydrous acctone, a white precipitate is obtained, which is filtered off and dried. The polymer contains 36.6% of Br. It is water-soluble.	50

5	EXAMPLE 22. Polymer of the formula I (with R=R'=CH ₂), A=(CH ₂), B=(CH ₂), and X=Br) A solution consisting of 130.2 g of N,N,N',N'-tetramethyl-trimethylenediamine and 216 g of 1,4-dibromobutane in 3,200 cm ³ of anhydrous methanol is heated under reflux for 55 hours. The polymer formed contains 46.2% of Br. It is soluble in water and in ethanol.	5
	EXAMPLE 23. Polymer of the formula I (with R=R'=CH ₂ , A=(CH ₂) ₆ , B=	
	$cH_2 \longrightarrow cH_2$ and $X = Br$)	
10	A mixture of 172.3 g of N,N,N',N'-tetramethyl-hexamethylenediamine and 264 g of p-xylylidene bromide in 3,200 cm ³ of anhydrous methanol is heated under reflux for 1 hour.	10
15	After cooling, the precipitate is filtered off and dried. The polymer contains 34.6% of Br. It is water-soluble.	15
	EXAMPLE 24. Polymer of the formula I (with R=R'=CH ₂ , A=(CH ₂) ₄ , B=(CH ₂) ₁₉ and X=Br) A solution consisting of 172.3 g of N,N,N',N'-tetramethyl-hexamethylenediamine	
20	and 300 g of 1,10-dibromodecane in 3,200 cm ³ of anhydrous methanol is heated under reflux for 15 hours. The polymer obtained contains 32.7% of Br. It is soluble in water and in ethanol.	20
25	EXAMPLE 25. Polymer of the formula I (with R=CH ₃ , R'=C ₄ H ₉ , A=(CH ₂) ₆ , B=(CH ₂) ₁₀ and X=Br) A solution of 214.4 g of N,N'-dibutyl-N,N'-dimethyl-trimethylenediamine and 300 g of 1,10-dibromodecane in 3,200 cm ³ of anhydrous methanol is heated under reflux for 70 hours. The polymer obtained contains 27.3% of Br.	25
	It is soluble in ethanol and in water-ethanol mixtures.	•
30	EXAMPLE 26. Polymer of the formula I (with R=R'=CH ₃ , A=(CH ₂) ₆ , B=(CH ₂) ₁₆ and X=Br) A solution consisting of 130.2 g of N,N,N',N'-tetramethyl-trimethylenediamine and 300 g of 1,10-dibromodecane in 3,200 cm ³ of anhydrous methanol is heated under	30
35	reflux for 38 hours. The polymer formed contains 34.3% of Br. It is soluble in water and in ethanol.	35
40	EXAMPLE 27. Polymer of the formula I (with R=CH ₃ , R'=C ₁₂ H ₂₅ , A=(CH ₂) ₆ , B=(CH ₂) ₄ and X=Br) A solution of 480.9 g of N,N'-didodecyi-N,N'-dimethyl-hexamethylenediamine and 216 g of 1,4-dibromobutane in a mixture of 2,000 cm ³ of acetonitrile and 4,000 cm ³ of isopropanol is heated for 12 hours at 85° C. The polymer formed contains 20.4% of Br. It is soluble in ethanoi.	- 40
45	EXAMPLE 28. Polymer of the formula I (with R=CH ₃ , R'=C ₁₂ H ₂₃ , A=(CH ₂) ₄ , B=(CH ₂) ₅ and X=Br) A solution of 480.9 g of N,N'-didodecyl-N,N'-dimethyl-hexamethylenediamine and 230 g of 1,5-dibromopentane in a mixture of 2,000 cm ³ of acetonitrile and 4,000 cm ³ of isopropanol is heated for 28 hours at 85° C. The polymer formed contains 19.9% of Br.	. 45
50	It is soluble in ethanol. On using methods analogous to those described in the preceding examples, the polymers of the formula I of which the structure is indicated in Table II below were obtained by Process 1 defined above.	. 50

			`			•										
I lb	Soluble in	water	ethanol, water-ethanol	ethanol, water-ethanol	ethanol	ethanol	ethanol	ethanol	ethanol	ethanol, water-ethanol	ethanol	ethanol	ethanol	water-ethanol, ethanol	water-ethanol, ethanol	water, water-ethanol,
-	. ×	Br	Br	Br	Br	Br	Br	, ä	Br	B	Br	Br	Br	Br	Br	Br
101	R'	£	C,1,H,18	C,4H,1	C,H,5	C,4H,4	C,H,	C,H,	C,H,	C,H,,	C,H,	с,н,	C,H,	cyclohexyl	iso-C,H,	CH,
1	R	CH,	CH.	CH,	Ж	СН,	СН,	CH,	. "НЭ	СН,	CH,	GH,	CH,	CH,	CH,	CH,
TABLE II	. B	(CH ₂),	(CH ₂),	(CH,),0	(CH ₂),	(CH ₂),	(CH ₂),	p-xylylene.	p-xylylene	(CH,),	(CH,),	(CH ₂),	(CH ₂),	(CH ₂),	(CH,),	a mix- (CH,),, 50 mol % ture of (CH,),0, 50 mol %
										•						
	٧	(CH ₂),	(CH ₂),	(CH ₂),	(CH ₂),	(CH ₂) ₃	(CH ₂),	(CH ₂),	(CH ₂),	(CH ₂),	(CH,),	(CH ₂),	(CH,),	(CH ₂),	(CH ₂),	(сн.),
	Ex.	53	30	31	32	~	34	35			38	39	_		42	£

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Hi.	Soluble in	water, water-ethanol	ethanol, water-ethanol	water, water-ethanol, ethanol	water, water-ethanol	water, water-ethanol	water, water-ethanol	ethanol	ethanol -	ethanol, water-ethanol	water, water-ethanol	water, water-ethanol, ethanol	water, water-ethanol, ethanol	water-ethanol ethanol
	×	Br	Br	Br	Br	Br	Br	Br	Ä	ğ	ğ.	Br.	ĕ	Br
	ж,	iso-C,H,	benzyl	°H,	CH,	CH,	CH,	C,H,,	С,Н",	C,H,	-(CH ₁) ₁ -0-(CH ₂) ₂ -	benzyl	с,́н,	iso-C,H,
itiiidatioii)	ĸ	CH,	CH,	сн,	СН,	сн,	CH,	CH,	Н	.	-(CH ₁) ₁ -	CH,	CH,	CH,
indianiiina ii centru	Ø	(CH,),	(CH ₂),	(CH,),	(CH,),	(CH,),	(CH ₂),	-7(СН')	p-xylylene	(CH ₂),	(CH,),	(CH,),	(CH2)	(CH,),
	A	(CH,),	(CH ₂)3	(CH ₁),	(CH ₂), +1.6% by weight of terminal groups -N(C,H ₆),	(CH ₂) ₄ + 2.8% by weight of terminal groups -N(C ₂ H ₂) ₁	(CH ₂) ₆ + 7.4% by weight of terminal groups -N(C ₂ H ₂),	(СН,),	(CH,),	. p-xylylidene	(CH ₂),	(CH ₂) ₆	m-xylylidene	(CH ₂),
	ŖŠ.	4	45	46	47.	84	49	20	51	25	53	54	55	S.

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	Soluble in	water, water-ethanol	water, water-ethanol, ethanol	ethanol	water, water-ethanol	water, water-ethanol	water, water-ethanol, ethanol	water, water-ethanol ethanol	water, water-ethanol, ethanol	water, water-ethanol, ethanol	ethanol	water, water-ethanol
,	×	Br	В	Br	Br		Br	Br	В	Br	Br	B.
	R',	С,Н,	c,H,	C,H,,	ĊH,	CH,	(CH ₂) ₂	. 	, _t	, tho	C,H,1	(CH,),-
tion)	æ	сн,	СН,	. CH³	. HO	CH,	-(CH ₁),-0-(CH ₁),-	. CH,	CH,	. .	ť	-(CH,-0-(CH,),-
TABLE II (continuation)	B	(CH ₃),	(CH;),	o-xylylene	o-xylylene	(CH,),	(CH,),	-сн,-сн,-сн- сн,	(CH ₂),	-(CH,),-OH	(GH,),	(CH ₂),
					-							
	A	(CH,),	(CH ₂),,	(CH,),	(CH,),	(CH,),	, (CH,),	(CH ₂),	(CH ₂), ₀	(CH ₂),	-Сн,-Сн,-Сн- Сн,	(CH ₂),
	R. No.	57	88	59	09	19	62	63	49	65	99	19

ontinuation)
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Ex.						
Š	٨	8	₩.	à	×	Soluble in
89	(CH ₃) _s	(Сн),	CH,	CH,	Br	water, water-ethanol, ethanol
69	(CH ₃),	(CH ₂),	CH,	ĊH,	В	water, water-ethanol
70	-CH ₂ -CH-(CH ₂) ₅ -CH-CH ₂ - C ₄ H ₅ C ₄ H ₅	(CH,),	СН		B	ethanol
71	-CH,-CH-(CH,),-CH-CH,- 	(Сн),	CH,	С,н,,	Br	ethanol
72	-cH,-CH-CH,- C,,H,,	(CH,),	CH,	СН,	, Br	water-ethano! ethano!
73	-CH,-CH-CH,- C,1H,s	(CH,),	СН	C,H,	Br	ethanol

EXAMPLE 74. Polymer of the formula I (with R=R'=CH₅, A=B=(CH₇)₁₀ and X=B₇)

50 g of 10-bromodecyl-dimethylamine hydrobromide, prepared according to M. R. LEHMAN, C. D. THOMPSON and C. S. MARVEL, J.A.C.S., 55, 1977 (1933) are dissolved in 200 cm² of water. An aqueous 25% sodium hydroxide solution is added until the pH is 12. The mixture is extracted with chloroform and the extracts are evaporated to dryness. The residue is dissolved in 250 cm² of methanol and the solution is heated under redux for 24 hours.

On addition of erival areaster.

On addition of ethyl acetate, a precipitate of the polymer formed, which contains 27.8% of Br, is obtained.

This is soluble in water and in ethanol and is practically identical with the

2

product described above in Example 9.

The 10-bromodecyl-dimethylamine hydrobromide used as the starting material is obtained as follows: 12.6 g of 10-phenoxydecyl-dimethylamine are dissolved in 63 cm³ of a 48% strength aqueous solution of hydrobromic acid. The solution is heated to 150° C, and is distilled under atmospheric pressure until the temperature of the vapours reaches 125° C, 63 cm³ of 48% strength hydrobromic acid are added and

and X=Br). Process 1a.

is heated under reflux for 60 hours.

55

A solution consisting of 160.3 g of 2,2'-bis-(dimethylamino)-diethyl ether, 264 g of para-xylylidene bromide, 1,000 cm³ of acetonitrile and 4,000 cm³ of isopropanol

The polymer obtained contains 33.8% of Br9. It is soluble in water and in a 50.50 water-chanol mixture. The quaternised polymers mentioned in the attached table were prepared analogously.

Ex. R. A B X Soluble in Process Process 82 CH,CH,OH CH, T (CH,J),								*;		-,									
R R' A B B' CH,CH,OH CH, CH, CH, CH, CH, CH,,	, (Process	1a	1a	. 1a		1a		:	. 1a		la	1	1a	1b	1a	1a	*	1a
R R' A B B CH,CH,OH CH, (CH,), (CH,), CH,. C		Soluble in	water water-ethanol	Water	Ethanol, water-ethanol		Ethanol	-		Ethanol, water-ethanol		P		Ethanol water-ethanol	Water, Water-ethanol	Water, water-ethanol	Ethanol, water-ethanol	Water, water-ethanol	•
# R R' A CH,CH,OH CH, ". CH,	• • • • • • • • • • • • • • • • • • •	×	Br	Br	Br	ĭ	Br	ă	i	Br	*	ğ		Br	Br	Br	Br	Br	Br
R R' R' CH,		B	(CH ₂),	(CH ₂),0	-сн,-снон-сн,-	. (CH ₂ CH ₂	(CH.)		•		_(CH ₂),		-(CH,),-0-(CH,),-	-сн,-снон-сн,	(CH,);		(CH ₂),	(CH ₂),
л сн, сн, он сн, сн, сн, сн, сн, сн, сн, сн, сн, сн	gously.	٧	(CH ₂),		•		•	(CH.)		(CH,),		-2#5-		(CH,),	(CH,),	-(CH ₂) ₂ -S-S-(CH ₂) ₂ -	•	2	
	analo	R,	сн	CH,	C,H,,		2	C.H.		CH,		CH,		C,H,	CH,	CH,	C,H,	· H	СН
R 88 88 88 88 89 89 89 89 89 89 89 89 89		~	сн,сн,он	:			. :			•		:			:	CH,	CH,	E	CH,
<u> </u>		Ex.	82	83	84		85	98		87		8		68	8	16	22	83	8

Process	1a	1a	1a	1a	e t	18	1a	1a	1a	15	91 9	d	19	19
Soluble in	Ethanol, water-ethanol	Water, water-ethanol	•	•	î	2	:	:	•	:	•	Water	Water, water-ethanol	4,
×	Br	Br	Br	Br	ğ	Br	ğ	B	Br	Br	Br	Br	Br	Br
В	(CH²)°	-CH2-CHOH-CH3-	(CH ₂),	-(CH ₂) ₂ -0-(CH ₂) ₂	-th	CH2—CH2	(CH ₂),	-(CH ₂) ₂ -0-(CH ₂) ₂ -	(CH,),	-сн,-снон-сн,-	-сн,-снон-сн,-	-сн,-снон-сн,	-сн,-снон-сн,	*:
¥	-(CH ₂) ₂ -S-S-(CH ₂) ₂ -	2					-(CH ₁),-S-S-(CH ₂),	$-(CH_2)_2-0-(CH_2)_2-$	$-(CH_2)_1-0-(CH_2)_1-$	-(CH ₂) ₃ -0-(CH ₂) ₂ -	(CH ₂),	(CH ₂),	(CH;),	(CH ₂),
							,							
R,	C ₁₁ H ₂₆	CH,	сн,	· HD	C,H,	, c	CH,	CH,	`#5	GH,	CH,	CH,	CH,	CH,
æ	CH,	, Æ	Ж	CH,	, ch,	CH,	CH,	Ж	CH,	CH,	CH,	CH,	Н	CH,
No.	95	%	16	86	66	100	101	102	103	104	105	106	107	108
	R R' A B X Soluble in	R R' A B X Soluble in CH ₃ CH ₃ , Br Ethanol, water-ethanol	R R' A B X Soluble in CH ₃ CH ₃ , CH ₃ , CH ₃ , CH ₃ , CH ₃ CH ₃ , CH ₃ , CH ₃ CH ₃ , CH ₃ CH ₃ , CH ₃	R R, A B X Soluble in CH, C,1,H,1 —(CH,2,1-S-S-(CH,2)2— (CH,3,2— CH,3,2— —(CH,2,2—CHOH—CH,2— Br Ethanol, water-ethanol CH, CH, ,, —(CH,2,1)2— Br Water, water-ethanol CH, CH, ,, (CH,3)40 Br Water, water-ethanol	R R A B X Soluble in CH, C,1,H,s —(CH,2,2-S-(CH,2)2- CCH,2,6 Br Ethanol, water-ethanol CH, CH, , —CH,2-CHOH—CH,2- Br Water, water-ethanol CH, CH, , (CH,1,0) Br " CH, CH, , —(CH,2,2-O-(CH,2)2 Br "	R R' A B X Soluble in CH3 CH4,2 -(CH2,2-S-CCH2,2- (CH3,6 Br Ethanol, water-ethanol CH3 CH3 " -CH2-CHOH-CH2- Br Water, water-ethanol CH3 CH3 " -CH2,0-CH3,0 Br " CH3 CH3 " -CH2,2-O-(CH2,2- Br " CH3 C,H3 " -CH2,2-O-(CH2,2- Br "	R R' A B X Soluble in CH3 CH3, G —(CH2, S—S—(CH2, 1)— —(CH2, 1)— B Ethanol, water-chlanol CH3 CH3 — CH2—CH0H—CH3— Br Water, water-chlanol CH3 CH3 — CH3, —0—(CH3, 1)— Br " CH4 CH3 — CH3, —0—(CH3, 2)— Br " CH4 CH5 — CH3, —0—(CH3, 2)— Br "	CH, R, A B X Soluble in CH, C,H,I,I -(CH,J,-S-S-(CH,J,-) (CH,J,I) Br Ethanol, water-othanol CH, CH,I -(CH,J,-S-S-(CH,J,-) Br Water, water-ethanol CH, CH,I -(CH,J,-O-(CH,J,-) Br CH, CH,I -(CH,J,-O-(CH,J,-) Br CH, CH, -(CH,J,-O-(CH,J,-) Br CH, CH, Br CH, CH,	CH, C, H, I A B X Soluble in CH, C, H, I −(CH,), -SS(CH,), - −(CH,), -SS(CH,), - Br Ethanol, water-ethanol CH, CH, −(CH,), -SS(CH,), - Br CH, CH, −(CH,), -D(CH,), - Br CH, CH, Br CH, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH,	R R? A B X Soluble in CH, 1 C,1,H,1 —(CH,1,-S-S-(CH,1,2) Br Ethanol, water-chianol CH, 2 CH, 3 —(CH,1,2-S-S-(CH,1,2) Br Water, water-chianol CH, 3 CH, 3 —(CH,1,2-O-(CH,1,2) Br CH, 3 CH, 3 —(CH,1,2-O-(CH,1,2) Br CH, 4 CH, 3 —(CH,1,2-O-(CH,1,2) Br CH, 4 CH, 4 —(CH,1,2-O-(CH,1,2) Br CH, 5 CH, 4 —(CH,1,2-O-(CH,1,2) Br CH, 5 —(CH,1,2-O-(CH,1,2) Br CH, 6 CH, 7 —(CH,1,2-O-(CH,1,2) Br	R R A B X Soluble in CH ₃ C ₁ ,H ₃ -(CH ₃ ,-S-S-(CH ₃)- (CH ₃),* Br Ethanol, water-chanol CH ₃ CH ₃ -(CH ₃ ,-CH ₃)- Br water, water-chanol CH ₃ CH ₃ -(CH ₃),-O-(CH ₃)- Br CH ₃ CH ₃ -(CH ₃),-O-(CH ₃)- Br CH ₄ CH ₃ -(CH ₃),-O-(CH ₃)- -(CH ₃),-O-(CH ₃)- Br CH ₄ CH ₃ -(CH ₃),-O-(CH ₃)- -(CH ₃),-O-(CH ₃)- Br CH ₄ CH ₃ -(CH ₃),-O-(CH ₃)- -(CH ₃),-O-(CH ₃)- Br CH ₄ CH ₄ -(CH ₄),-O-(CH ₃)- -(CH ₄),-O-(CH ₃)- Br	CH, R, A B X Soluble in CH, C,1,H ₁ —(CH,1,-S-S-(CH,1,- Br Ethanol, water-ethanol CH, CH, — (CH,1,-S-S-(CH,1,- Br Water, water-ethanol CH, CH, — (CH,1,-CHOH—CH,1- Br " CH, CH, — (CH,1,-CHOH—CH,1- Br " CH, CH, — (CH,1,-CHOH—CH,1- Br " CH, CH, — (CH,1,-CHOH,1- Br " CH, CH, — (CH,1,-CHOH,1- Br " CH, CH, — (CH,1,-CHOH,1- Br " CH, CH, — (CH,1,-CH,1,1- — (CH,1,1- Br " CH, CH, — (CH,1,-CH,1,1- — (CH,1,1- Br " CH, CH, — (CH,1,-CH,1,1- — (CH,1,1- Br " CH, CH, — (CH,1,1- — (CH,1,1- Br " CH, CH, — (CH,1,1- — (CH, C, H, I — (CH, J, -S-C(H, J, -CH, J, -S-C(H, J, -CH, J, -S-C(H, J, -CH, -CH, -CH, J, -CH, -CH, -CH, J, -CH, -CH, J, -CH, -CH, J, -CH, -CH, -CH, -CH, J, -CH, -CH, -CH, J, -CH, -CH, J, -CH, -CH, J, -CH, -CH, -CH, -CH, -CH, J, -CH, -CH, -CH, -CH, -CH, -CH, -CH, -CH	CH, R, A B X Soluble in CH, CH, —(CH,),~S-S-(CH,)~ —(CH,),* Br Ethanol, water-chanol CH, CH, —(CH,) —(CH,),* Br —(CH,), water-chanol CH, CH, —(CH,) —(CH,),* Br ————————————————————————————————————

Process	92	16		16	la	1a	Ia	1a	İa	la -	Ta	1a	1a	18	1a
Soluble in	Water, ethanol	Water, ethanol		Water, water-ethanol	:	**			Water, ethanol	Water, water-ethanol	Ethanol	Water, water-ethanol	Ethanol, water-ethanol	Water, ethanol	44
"×	Ä	Ŗ		Ä	Ŗ	Br	Br	Br	-Br	ğ	ž.	ij	B	Br	Br
В	-сн,-снон-сн,	-сн,-снон-сн,			(CH,),	(CH ₂),	(CH ₂),	-сн'-снон-сн'-	СН,СНОНСН,	-(CH ₂),-0-(CH ₂),-	•	an		(CH,),	(CH ₁),
V	(CH ₂),	(CH,),		CH2 CH1	-(CH ₂) ₁ -S-(CH ₂) ₂ -	-(CH ₂) ₂ -SO-(CH ₂) ₂ -	-(CH ₂) ₁ SO ₁ -(CH ₂) ₁	(СН,),	(CH ₂), ₁₀	CH ₂ —Ch ₂	(Сн,),	(CH,),	(CH,),	(CH,),	(CH ₂),
	-(CH ₂),	CH.		.#5	£	Сн.	. #J	CH,	C,II,	"H"	C,,H,,	CH,	· C,H,,	СН	CH,
æ		CH,		CH,	· H	CH,	сн,	.	CH,	. CH,	Ť	CH,	CH,	Ж	сн,
Z. So.	109	110		1111	112	113	114	1115	116	117	118	119	120	121	122

26					1,515,072			
	Process	1a	la ,	1a	n T	. La	1.6	15
	Soluble in	Water	Water, ethanol	Water	Alcohol	Water, ethanol	Water	Water
	x	Br	B	. Br	Br	Br	В	Br
	В	CH2 CH2	—(CH ₂),—CH—	(CH ₂),	-CH,CHCH, 0H	(CH ₁) ₂ -0-(CH ₂) ₂	сн, —сн —сн, Он	сн, сн сн, ОН
	А	'('HO)	(CH ₂) ₅	(CH ₂),-0-(CH ₁),	CH2-CH2-CH2-CH2-CH2-CH2-CH2-CH2-CH2-CH2-	-C+10-C→	-(CH ₁) ₁ -CH- CH ₁	
	R,	CH,	CH,	сн,	CH,	Ŧ.	. 19	·
					34		_(CH ₃),	-(CH ₂),
	æ	. H	Ж	Ë	.	. 		1
· .	N. S.	123	124	125	126	127	128	129

Process	la	. ra	1a		e.		1a	1a	Ta
Soluble in	Water, ethanol	Water	Water		Water, ethanol	Water, ethanol	Ethanol	Water	Water, ethanol
×	Br	Br	Br	·.	. Br	Br	ō .	В	Br
В	(CH ₃),	(CH ₂),	-(CH ₂) ₃ -CH-	CH,	CH2 CH2	Сн,—сн.—сн,	сн,—сн—сн, он	CH,-CH-CH,	Сн,—Сн.—Сн, Он
Y	(CH ₂),	(CH ₂),	(CH ₂),		(CH,),	(CH,),	(CH ₂),	££	CH2———CH2
R,	iso C,H,	iso C,H,	C,H,		C,H,	ben zy l		C,H,	CH,
R	сн,	CH,	CH,		CH.	CH,	ĞH,	ch,	
RX.	130	131	132		133	134	135	136	137

ł Hp.	Process	la 1	la La				
1	Soluble in	Water, ethanol	2 2	S	10	a to 15 1ple 15	
*	×	Br	Br Br	SMETIC		use the crean rmer of Exan	
	В	(CH ₂),0	(CH;),	EXAMPLES OF COSMETIC COMPOSITIONS AND OF COSMETIC TREATMENT EXAMPLE I. Hand treatment creams. The following cream is prepared:	10 g 6 g 6 g 2 g 001 g 100 g	This cream is applied to the hands, whilst nubbing them to cause the cream to penetrate. The hands are soft and pleasant to the touch. 2. Analogous results are obtained by replacing the 4 grams of polymer of Example 1 in the above cream formulation by 3.5 grams of polymer of Example 15.	-
	V	CH-7	(CH ₂),	EXAMPLES OF COSMETIC CONTREASES TREASES IN The following cream is prepared:	"Vaseline" oil Cetyl alcohol Self-emulsifiable glyceryl monostearate Triethanolamine Methyl p-hydroxybenzoate Polymer of Example 1 Water, q.s.p.	This cream is applied to the hands, whilst trate. The hands are soft and pleasant to the touch. 2. Analogous results are obtained by replacir the above cream formulation by 3.5 grams	
				EXA.		This penetrate. The h	
•	à	C,H,	C,11,	5	10	pen 15	
	æ	CH,	CH,	* • .			
		•					

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	EXAMPLE II. Carrier creams for dyeing (oxidation dyeing). 1. A cream of the following formulation is prepared:	·
	Cetyl/stearyl alcohol 20 g	
	Oleyl diethanolamide 4 g	
5	Sodium cetyl/stearyl-sulphate 3 g	5
	Polymer of Example 15 5 g	
	Ammonia, 22°B strength (11N) 10 ml	
	m-Diaminoanisole sulphate 0.048 g	
	Resorcinol 0.420 g	
10	m-Aminophenol base 0.150 g	10
	Nitro-p-phenylenediamine 0.085 g	
	p-Toluylenediamine 0.004 g	
	Triion B(g) 1 g	
	Sodium bisulphite, d=1.32 1,200 g	15
15	Water, q.s.p. 100 g	15
	© "Trilon" B: the tetrasodium salt of ethylenediaminetetra-acetic acid. ["Trilon" is a Registered Trade Mark.]	
	30 g of this cream are mixed with 45 g of hydrogen peroxide of 20 volumes	
	strength. A smooth thick cream which is pleasant to apply and adheres well to the hair	
20	is obtained.	20
20 .	After waiting for 30 minutes, the hair is rinsed and dried.	
	On 100% white hair, a blond shade is obtained. The hair is easy to comb out	
	whether wet or dry. The hair looks shiny and is pleasant and silky to the touch.	
	2. A cream of the following formulation is prepared:	
25	Cetyl/stearyl alcohol 20 g	. 25
	Oleyl diethanolamide 4 g	
	Sodium cetyl/stearyl-sulphate 3 g	
	Polymer of Example 3 5 g	
	Ammonia, 22°Be strength (11N) 12 ml	
30	m-Diaminoanisole sulphate 0.048 g	30
	Resorcinol 0.420 g	
	m-Aminophenol base 0.150 g	•
	Nitro-p-phenylenediamine 0.085 g	
	p-Toluylenediamine 0.004 g	25
35	Trilon B 1 g	35
	Sodium bisulphite, d=1.32 1,200 g	
	Water, q.s.p. 100 g	
	30 g of this cream are mixed with 45 g of hydrogen peroxide of 20 volumes	
	strength. A smooth thick cream which is pleasant to apply and adheres well to the	•
40	hair is obtained.	40
	After waiting for 30 minutes, the hair is rinsed and dried.	
	On 100% white hair, a blond shade is obtained. The hair is easy to comb out	
	either wet or dry. The hair looks shiny and is pleasant and silky to the touch.	
	EXAMPLE III. Wavesetting lotion for sensitised hair.	46
45	1. An alcoholic lotion of the following formulation is prepared:	45
	Polanda de la companya del companya de la companya del companya de la companya de	
	Polyvinylpyrrolidone 1 g	
	Polymer of Example 7 1 g	
	Ethyl alcohol, q.s.p. 100 ml	
	This solution is applied to the hair, the hair is set in waves and is dried.	
50	The hair is strengthened and plasticised.	50
50	The hair is shiny and bulky; it is silky to the touch and easy to comb out.	50
	2. Analogous results are obtained on replacing the polymer of Example 7 in the	
	above formulation by the polymer of Example 16.	
	3. The lotion of the following formulation is prepared:	
55	Polymer of Example 18 0.8 g	55
	Polyvinylpyrrolidone/vinyl acetate, 60:40 1.0 g	
•	Triethanolamine, q.s. pH 6	
	Water, q.s.p. 100 ml	

	•				1 1	
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30		1,513,672				30
,•		The lotion is applied to bleached hair. The hair is set in Analogous results to those of the preceding example are obta 4. The lotion of the following formulation is prepared:		and dried.		
5		Polymer of Example 19 Polyvinylpyrrolidone/vinyl acetate, 60:40 copolymer Ethyl alcohol, q.s. 50° Triethanolamine, q.s. pH 7	7	g g	4	5
		Water, q.s.p.	100	ml	1.	
10		The lotion is applied to bleached hair. The hair is set in wav Analogous results to those of the preceding example are obta 5. The lotion of the following formulation is prepared:		dried.	·.	10
15	. !	Polymer of Example 17 Vinyl acetate/crotonic acid, 90:10 copolymer Triethanolamine, q.s. pH 7.5 Water, q.s.p.		S g S g ml		. 15
		The lotion is applied to bleached hair. The hair is set in wav Analogous results to those of the preceding example are obta 8. The following solution is prepared:		dried.		
20		Polymer of Example 2 Vinyl acetate/crotonic acid, 90:10 copolymer Monoethanolamine, q.s. pH 7 Water, q.s.p.		5 g 5 g ml		20
25		The solution is applied to bleached hair. The hair is set an Analogous results to those of the preceding example are obtained as a set of the preceding example are obtained as a set of the following solution are applied to clean, wet have been applied to clean, wet have the set of the following solution are applied to clean, wet have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the set of the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, we have the following solution are applied to clean, and the following solution are applied	ined. ith rin	sing)	. •	25
		Monoethanolamine, q.s. pH 7.5 Water, q.s.p.	100	ml		
30		The lotion is allowed to act for 5 minutes and the hair is the The hair is soft to the touch and easy to comb out. It is set in waves and dried. The dry hair is easy to comb out.	n rinse	ed.		30
35		The hair is shiny and springy, and has body. 2. 25 ml of the following solution are applied to clean, wet h	аіг:	÷.		35
		Polymer of Example 12 Citric acid, q.s. pH 6	6	g		
40		Water, q.s.p. The lotion is allowed to act for 5 minutes and the hair if the hair is soft to the touch and easy to comb out. It is set in waves and dried. The dry hair is easy to comb out. The hair is shiny and springy, and has body.	100 s then	ml rinsed.		40
		3. 25 ml of the following solution are applied to clean, wet h	air:			
45		Polymer of Example 21 Triethanolamine, q.s. pH 6 Water, q.s.p.	6 100	g ml	· .	45
50		The lotion is allowed to act for 5 minutes and the hair is the The hair is soft to the touch and easy to comb out. It is set in waves and dried. The dry hair is easy to comb out.	n rinse	ed.		50

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•	_31	1,513,672		31	
•		The hair is shiny and springy, and has body. 4. 30 ml of the following solution are applied to clean, we	et hair:		•
. '	. 5	Polymer of Example 22 Water, q.s.p. The pH is about 7.	7 g 100 ml	5	•
* *	3		•		
		The lotion is allowed to act for 5 minutes and the hair is The hair is soft to the touch and easy to comb out. It is set in waves and dried. The dry hair is easy to comb out.	then rinsed.	1	
	10	The hair is shiny and springy, and has body. 5. 25 ml of the following solution are applied to clean, w	et hair:	10	
		Polymer of Example 23 Monoethanolamine, q.s. pH 5	5 g	•	•
		Water, q.s.p.	100 ml		
	15	The lotion is allowed to act for 5 minutes and the hair is The hair is soft to the touch and easy to comb out. It is set in waves and dried. The dry hair is easy to comb out.	then rinsed.	15	
		The hair is shiny and springy, and has body.	•		• .
	20	EXAMPLE V.—Structuring lotion (applied wind) 1. Prior to use, 0.3 g of N,N'-di-hydroxymethylethylen to as compound A, is mixed with 25 ml of a solution containing	ethiourea, hereafter referred	20	• .
		Polymer of Example 11 Hydrochloric acid, q.s. pH 2.7	0.4 g	•	
	25	Water, q.s.p.	100 ml	25	
		The mixture is applied to hair which has been washe setting it in waves. The hair can be combed out easily and is silky to the tou It is set in waves and dried.			
	30	The hair is shiny and springy; it has body (bulk), i easy to comb out. 2. Prior to use, 0.4 g of compound A is mixed with 2		30	
		ing:	•	•	
	35	Polymer of Example 14 Phosphoric acid, q.s. pH 2.7 Water, q.s.p.	0.5 g 100 ml	35	. * •
		The mixture is applied to hair which has been washe setting it in waves.	·		
	40	The hair can be combed out easily and is silky to the touc It is set in waves and dried. The hair is shiny and springy; it has body (bulk), is easy to comb out.		40	
•		3. Prior to use, 0.5 g of compound A is mixed with 2 ing:	5 ml of a solution contain-		
÷ •	45	Polymer of Example 24 Phosphoric acid, q.s. pH 3	0.6 g	45	
		Water, q.s.p.	100 mi		•
		The mixture is applied to hair which has been washes setting it in waves.			
·	50	The hair can be combed out easily and is silky to the touc It is set in waves and dried.		50	
	•	The hair is shiny and springy; it has body (bulk), is easy to comb out.	silky to the touch and is		

- 33	1,072	33
,	4. Prior to use, 2 g of compound A are mixed with 25 ml of a solution containing:	
5	Polymer of Example 1 5 g Phosphoric acid, q.s. pH 2.8 Water, q.s.p. 100 ml	5
10	The mixture is applied to hair which has been washed and towelled dry. It is left to act for 10 minutes, and the hair is rinsed. The hair can be combed out easily and is soft (silky) to the touch. It is set in waves and dried under a hood. The dry hair can be combed out easily. It is shiny and springy and has body (bulk). 5. Prior to use, 1.5 g of compound A are mixed with 25 ml of a solution con-	10
15	Polymer of Example 11 5.5 g Phosphoric acid, q.s. pH 3 Water, q.s.p. 100 ml	15
20	The mixture is applied to hair which has been washed and towelled dry. It is left to act for 10 minutes, and the hair is rinsed. The hair can be combed out easily and is soft (siky) to the touch. It is set in waves and dried under a hood. The dry hair can be combed out easily. It is shiny and springy and has body (bulk).	20
	EXAMPLE VII—Shampoos. 1. The following solution is prepared:	
25	C ₁₁ —C ₁₄ \(\alpha\)-diol condensed with 3 to 4 molecules of glycidol Polymer of Example 1 Lactic acid, q.s.p. Water, q.s.p. 100 cm ³	25
30	When applied to the head, this solution, which has a limpid appearance, produces a copious and rather soft foam and assists the combing out of the wet hair. After drying, the hair is springy, light and shiny. 2. The following solution is prepared:	30
35	C ₁₂ H ₂₂ —(O—CH ₂ —CHOH—CH ₂ —) OH (lauryl alcohol condensed with 4 molecules of glycidol or epichlorohydrin) Polymer of Example 1 Terriory of Example 1 Terriory of Example 1 Terriory of Example 1	35
40	Tertiary stearylamine polyoxyethylenated with 5 mols of ethylene oxide 1.5 g Lactic acid, q.s.p. Water, q.s.p. 100 cm ³	40
45	When applied to the head, this solution, of limpid appearance, produces a copious and soft foam which is readily removed by rinsing. The hair is very easy to comb out and after drying possesses bulk and springiness whilst remaining soft and easy to style. 3. The following solution is prepared:	45
50	C ₁₁ —C ₁₄ a-diol condensed with 3 to 4 molecules of glycidol 17 g Polymer of Example 12 3 g Lactic acid, q.s.p. pH 3.5 Water, q.s.p. 100 cm ³	50
	When applied to the head, this solution, of limpid appearance, produces a	

When applied to the head, this solution, of limpid appearance, produces a copious and rather soft foam and makes it possible to improve the combing out of the wet hair. After drying, the hair is soft and shiny and has a lightweight appearance.

35	1,072,072	33
	either wet or dry. The hair has a shiny appearance and is pleasant and silky to the touch.	
	The same result is obtained on replacing the compound of Example 105 by one of the compounds of the following examples:	•
5	Example 106 5%	5
	Example 107 5%	
	Example 108 6%	
	Example 110 4.5% Example 111 6%	
10	Example 111 6% Example 76 3%	10
	2. A cream of the following formulation is prepared:	
	Stearyl alcohol 18 g Coconut monoethanolamide 6 g	
	American In the 1 to 1000/ 56 to 1 to 10 to 10	
15	Command of Emanda 110	15
	Ammonia, 22° Bé strength (11 N) 10 cm ³	13
	m-Diaminoanisole sulphate 0.048 g	
	Resorcinol 0.420 g	
	m-Aminophenol base 0.150 g	
20	Nitro p-phenylenediamine 0.085 g	20
	p-Toluylenediamine 0.004 g	
	Trilon B Sodium bisulphite (d=1.32) 1.000 g 1.200 g	
	Wassa	
	water, q.s.p. 100 g	
25	30 g of this formulation are mixed with 45 g of hydrogen peroxide of 20 volumes	
	strength. A smooth, thick cream is obtained, which is pleasant to apply and which	
	adheres well to the hair.	
	After waiting for 30 minutes, the hair is rinsed and dried.	
20	On 100% white hair, a blond shade is obtained. The hair is easy to comb out	
30	either wet or dry. The hair has a shiny appearance and is pleasant and silky to the touch.	1 .
	The same result is obtained on replacing the compound of Example 119 by one	
	of the compounds of the following examples:	
	• • • • • • • • • • • • • • • • • • •	
	Example 75 5%	
35	Example 104 4%	35
	Example 102 5%	
	Example 81 5.5%	
	Example 103 6%	
	EXAMPLE XI.—DYEING SHAMPOOS.	
40	1. A dyeing shampoo of the following formulation is prepared:	40
	Nonylphenol+4 mols of ethylene oxide 25 g	
	Nonyiphenol + 9 mols of ethylene oxide 23 g	
	Compound of Example 89	
40	Ethyl alcohol, 96% strength 7	
45	rropylene glycol 14 6	45
	Ammonia, 22 Be strength (11 N) 10 cm ³	
	m-Diaminoanisole sulphate 0.030 g	
	Resorcinol 0.400 g	
50	m-Aminophenol base 0.150 g	(2)
	p-Aminophenol base 0.087 g Nitro p-phenylenediamine 1.000 g	50
	0 1) 44 44 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	
	W/	
55	50 g of this formulation are mixed with the same quantity of hydrogen peroxide	55
	of 20 volumes strength and the gel obtained is applied to hair by means of a brigh	55
	The mixture is left for 30 minutes and the hair is then rinsed.	

			
	The hair is easy to brush out and is silky to dried.	the touch. It is set in waves and	
•	The hair is shiny and springy and has body	(bulk); it is silky to the touch and	
5	easy to comb out. A chestnut tint is obtained on a brown backgrou 2. A dyeing shampoo of the following formula		5
	Nonylphenol+4 mols of ethylene oxide	25 g	
	Nonylphenol+9 mols of ethylene oxide	23 g	
10	Compound of Example 118 Ethyl alcohol, 96% strength	5 g r.	10
	Propylene glycol	7 g 14 g	
	Ammonia, 22 Bé strength (11 N)	10 cm ^s	
	m-Diaminoanisole sulphate	0.030 g	
15 -	Resorcinol m-Aminophenol base	0.400 g 0.150 g	15
	p-Aminophenol base	0.087 g	
	Nitro p-phenylenediamine	1.000 g	
	Trilon B	3.000 g	
20	Sodium bisulphite (d=1.32)	1.200 g	20
٠.	Water, q.s.p.	100 g	
	50 g of this formulation are mixed with the solution of 20 volumes strength and the gel obtained is an The mixture is left for 30 minutes and the hair The hair is easy to brush out and is silky to	plied to hair by means of a brush. is then rinsed.	
25	dried.	the touch. It is set in waves and	25
	The hair is shiny and springy and has body	(bulk); it is silky to the touch and	
	easy to comb out. A chestnut tint is obtained on a brown backgrou	nd.	
30	EXAMPLE XII.—TREATMENT LOTION 30 ml of the following solution are applied to cl	N (application with rinsing). ean, wet hair:	30
	Compound of Example 94	5 g	
	Monoethanolamine, q.s.	pH 7.5	
	Water, q.s.p.	100 cm ³	
	After waiting for 5 minutes, the hair is rinsed.		
35	The hair is soft to the touch and can be combed	out easily.	35
	It is set in waves and dried.	•	•
	The dry hair can be combed out easily. It is The same result is obtained by replacing the	shiny and springy and has body.	
	of the compounds of the following examples:	compound of Example 94 by one	
40	Example 91	4 g	40
		4 g 6 g	
	Example 93	6 g	
	Example 100 Example 97	4 g	
15	Example 112	6 g 5 g	45
	Example 96	5 g	45
	Example 98	6.5 g	
	Example 101 Example 99	4.5 g	
	•	5 g	
60	EXAMPLE XIII.—STRUCTURING LOTIC 1. Prior to use, 2 g of N,N'-di-hydroxymeth 25 cm ⁵ of a solution containing:	ON (application with rinsing). Sylethylenethiourea are mixed with	50
	Compound of Example 117	5 -	
	Hydrochloric acid, q.s.	5 g pH 2.5 100 cm ³	
55	Water, q.s.p.	100 cm ³	55
	The mixture is applied to hair which has been		
	The mixture is applied to hair which has been w	asned and towelled dry.	

37	1,513,672	37
	After waiting for 10 minutes the hair is rinsed. The hair is easy to comb out and is soft (silky) to the touch. It is set in waves and dried under a hood.	
5	The dry hair can be combed out easily, is shiny and springy and has body (bulk). 2. Prior to use, 2 g of N,N'-di-hydroxymethylethylenethiourea are mixed with 25 cm ³ of a solution containing:	
10	Compound of Example 79 Hydrochloric acid, q.s. Water, q.s.p. 3 pH 2.5 Water, q.s.p. 100 cm ³	10
15	The mixture is applied to hair which has been washed and towelled dry. After waiting for 10 minutes the hair is rinsed. The hair is easy to comb out and is soft (silky) to the touch. It is set in waves and dried under a hood. The dry hair can be combed out easily, is shiny and springy and has body (bulk).	15
	3. Prior to use, 2 g of N,N'-di-hydroxymethylethylenethiourea are mixed with 25 cm ² of a solution containing:	
20	Compound of Example 109 4 g Hydrochloric acid, q.s. pH 2.5 Water, q.s.p. 100 cm ⁵	20
25	The mixture is applied to hair which has been washed and towelled dry. After waiting for 10 minutes the hair is rinsed. The hair is easy to comb out and is soft (silky) to the touch. It is set in waves and dried under a hood. The dry hair can be combed out easily, is shiny and springy and has body (bulk).	25
 30	EXAMPLE XIV.—STRUCTURING LOTION (application without rinsing). Prior to use, 0.3 g of N,N'-di-hydroxymethylethylenethiourea is mixed with 25 cm ³ of a solution containing:	30
	Compound of Example 82 Phosphoric acid, q.s. Water, q.s.p. 0.5 g pH 2.8 100 cm ³	
35	The mixture is applied to hair which has been washed and dried, before setting it in waves. The hair is easy to comb out and is silky to the touch. It is set in waves and dried.	35
40	The hair is shiny and springy, it has body (bulk), is silky to the touch and is easy to comb out. The same result is obtained on replacing the compound of Example 82 by one of the compounds of the following examples:	40
45	Example 77 0.4 g Example 87 0.6 g Example 83 0.4 g	
	Example 88 0.5 g EXAMPLE XV.—WAVESETTING LOTION FOR SENSITISED HAIR. 1. A wavesetting lotion of the following formulation is prepared:	45
50	Polyvinylpyrrolidone Compound of Example 92 Ethyl alcohol, q.s. 1 g 2.5 g 100 cm ^s	50
	The solution is applied to the hair. The hair is set in waves and dried. The hair is strengthened and plasticised. It is shiny and bulky: it is silky to the touch and are the same and the same and the same are the same and the same are t	

The hair is strengthened and plasticised.

It is shiny and bulky; it is silky to the touch and easy to comb out.

After setting in waves and drying, the hair combs out easily and is soft to the

The same solution can be packaged as an aerosol with nitrogen or nitrous oxide

45

The hair is shiny and springy.

or freons as propellants.

,	1.512.773	20.
39	1,513,672	39
•	EXAMPLE XVIII.—TREATMENT LOTION FOR USE BEFORE DYEING. 20 cm ³ of the following solution are applied to dry, dirty hair:	•
•	Compound of Example 1 3 g	
	Monoethanolamine q.s.p. pH 8	
5	Water, q.s.p. 100 g	5
	After waiting for 5 minutes, a conventional ammoniacal oxidation dye is applied. It is left for 30 minutes.	
	After rinsing and giving an anionic shampoo, the hair combs out very easily. After wavesetting and drying, the hair is silky, shiny, springy and easy to style.	
10	EXAMPLE XIX—ANIONIC SHAMPOO.	10
	Triethanolamine lauryl-sulphate 10 g	
	Compound of Example 17 1 g Triethanolamine, q.s.p. pH 8	
	Triethanolamine, q.s.p. pH 8 Water, q.s.p. 100 g	
15		
13	EXAMPLE XX.—PRE-SHAMPOOING COMPOSITION.	15
,	Compound of Example 17 2 g	
•	Monoethanolamine, q.s.p. pH 7	
	Water, q.s.p. 100 g	
20	10 g of this composition are applied to dry, dirty hair. After waiting for 2 minutes, the hair is given a conventional anionic shampoo. The hair is easy to comb out and is soft to the touch both when wet and after	20
	drying. Analogous results are obtained on replacing the compound of Example 17 by the	
	compounds of Examples 1, 7, 13, 20, 22, 23, 46, 75, 76, 91, 92, 93, 105, 108, 110,	
25	111, 112, 117, 123, 130, 136 or 140.	25
	Analogous pre-shampooing compositions were prepared in the form of aerosol packs, using the same compounds. For example, the following procedure can be followed:	1
1	The following solution is prepared:	
30	Compound of Example 17 8 g	30
	Monoethanolamine, q.s.p. pH 7	
	Water, q.s.p. 100 g	
	25 g of this solution are introduced into an aerosol container and nitrogen is then introduced until the pressure reaches 12 kg/cm ² .	
35	Dry hair which is to be washed is impregnated using the aerosol pack thus obtained and after waiting for a few minutes the hair is given a conventional anionic shampoo.	35
	EXAMPLE XXI.—PRE-SHAMPOOING COMPOSITION IN THE FORM OF	
40	AN AEROSOL FOAM.	
40	The following formulation is prepared as an aerosol:	.40
	Na cetyl/stearyl-sulphate 1.3 g Nonylphenol ethoxylated with 4 mols of ethylene oxide 2.5 g Nonylphenol ethoxylated with 9 mols of ethylene oxide 1.5 g	
A.E.	Compound of Example 17 3.0 g	
45	Water 81.7 g "Freon" 114—"Freon" 12 mixture (70:30) 10.0 g Degree of filling 65%.	. 45
ΕΛ	The foam is applied to dirty, dry hair whilst rubbing so as to cause the product to penetrate thoroughly into the hair.	
50	A conventional anionic shampoo is applied. After waiting for 2 to 3 minutes, the hair is rinsed. The hair is soft to the touch and is easy to comb out.	50
	The hair is set in waves and dried. The dry hair combs out easily. It is shiny and springy and has body (bulk).	

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It should be noted that some of the polymers used in the present invention are described and claimed in our Application No. 20683/75 (Serial No. 1,513,671), which provides a quaternised polymer having recurring units of the formula:

a)
$$- \begin{bmatrix} \frac{CH_3}{I_1} \frac{A_{\square}}{A_1} - \frac{CH_3}{I_2} \frac{A_{\square}}{B_1} \\ \frac{I_1}{B_2} - \frac{I_1}{B_2} \end{bmatrix}$$
 (IIA)

in which each A₁ radical, which may be the same or different, represents either A, A being a group of the formula:

in which x, y and t each independently is 0 or an integer from 1 to 11, and E and K each independently is a hydrogen atom or an aliphatic radical, such that the radical contains less than 18 carbon atoms, or

A₁ represents a group of the formula:

$$\begin{array}{lll} --(CH_2)_n-O-(CH_2)_n-, \\ --(CH_2)_n-S-(CH_2)_n-, \\ --(CH_2)_n-S-S-(CH_2)_n-, \\ --(CH_2)_n-SO-(CH_2)_n-, \\ --(CH_2)_n-SO_2-(CH_2)_n- \end{array}$$

in which n is 2 or 3; X^{\ominus} is an anion derived from an inorganic or organic acid; R'_2 is an alkyl, alicyclic or arylaliphatic radical containing at most 20 carbon atoms; each B_1 radical, which may be the same or different, represents either B, B being a group of the formula:

in which D and G each independently is a hydrogen atom or an aliphatic radical, and v, z and u each independently is 0 or an integer from 1 to 11, such that up to two of them can be 0 and such that the radical contains less than 18 carbon atoms and the sum (v+z+u) is greater than 1 when the sum (x+y+t) is equal to zero, or B_1 represents a group of the formula:

with the proviso that R'_2 has at least 3 carbon atoms when A_1 or B_1 represents $-(CH_2)_n$ —O— $(CH_2)_n$ —, or when, simultaneously, A_1 =A and B_1 =B and that R'_2 does not represent a methyl radical when A_1 =A or A_1 represents

$$b) \qquad \frac{-(CH_{2})_{n}-O-(CH_{2})_{n}-or-(CH_{2})-S-(CH_{2})_{n}-or-(CH_{2})_{$$

in which each A_2 radical, which may be the same or different, represents either A as defined above, or a group of the formula:

$$\begin{array}{c} -(CH_2)_n - O - (CH_2)_n -, \\ -(CH_2)_n - S - (CH_2)_n -, \\ -(CH_2)_n - S - S - (CH_2)_n -, \\ -(CH_2)_n - S O - (CH_2)_n -, \\ - CH_2 - CH_2 - CH_2 -, \end{array}$$

n, X, R'2 and B1 being defined as above; or

c)
$$\frac{1}{\sqrt{2}} \frac{\chi^{\Theta}}{\sqrt{2}} \Lambda_{3} \frac{\chi^{\Theta}}{\sqrt{2}} CH_{2} - CH_{2} - CH_{2}$$
 (IIC)

in which Z represents an oxygen atom or the —CH₂— group; each A, radical, which may be the same or different, represents either A as defined above, or

10

$$-(CH_2)_n-O-(CH_2)_n-$$

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and X and n are as defined above as well as compositions containing such a polymer and a method of treating the skin or hair with it. No claim is made herein to the use of these polymers.

Subject to this disclaimer: WHAT WE CLAIM IS:-

15

1. A method of treating the hair and/or skin which comprises applying thereto at least one quaternised polymer possessing recurring units of the general formula:

15.

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in which: each R radical, which may be the same or different, is a C₁ to C₅ alkyl group or a —CH₂—CH₂OH group, each R' radical, which may be the same or different, is an aliphatic radical, an alicyclic radical or an araliphatic radical, which radical contains at most 20 carbon atoms, or R and R' attached to the same nitrogen atom form, with the latter, a ring which can contain a second hetero-atom other than nitrogen, each A radical, which may be the same or different, represents a divalent group having one of the formulae:

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25

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(ii)
$$-(CH_2)_y$$
- CH - $(CH_2)_z$ - CH - $(CH_2)_t$ - K

in which x, y and t are each independently 0 or an integer from 1 to 11 such that the sum (x+y+t) is 0 to 17, and E and K are each independently a hydrogen atom or an aliphatic radical having fewer than 18 carbon atoms,

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in which n is 2 or 3, each B radical, which may be the same or different, represents a divalent group having one of the formulae:

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(i)
$$\begin{array}{ccc} -\mathbb{C}H_2 - & & \\ & & & \\ \mathbb{C}H_2 - & & \\ & & & \\$$

in which D and G each independently are a hydrogen atom or an aliphatic radical having less than 18 carbon atoms, and v, z and u are each independently 0 or an integer from 1 to 11, and any two of them can simultaneously be 0, such that the sum (v+z+u) is at least 1 but not greater than 17 and the sum (v+z+u) is greater than 1 if the sum (x+y+t) is 0,

n being defined as above, and X² represents an anion derived from an organic or inorganic acid.

2. A method according to claim 1, in which A represents an o-, m- or p-xylylene group or a group of the formula:

in which E, K, x, y and t are defined as in claim 1, and B represents an o-, m- or p-xylylene group or a group of the formula:

in which D, G, v, z and u are defined as in claim 1.

3. A method according to claim 1 or 2, in which R is a methyl or hydroxyethyl radical, R' is an alkyl radical having 1 to 16 carbon atoms, a benzyl radical or a cyclohexyl radical, or R and R' together represent a polymethylene radical having 2 to 6 carbon atoms or the —(CH₂)₂—O—(CH₂)₂— radical; A is a xylyiene radical, a polymethylene radical having 2 to 12 carbon atoms which is optionally substituted by one or two alkyl radicals having 1 to 12 carbon atoms, or a radical of the formula:

25

—(CH₂)_n—S—(CH₂)_n—, —(CH₂)_n—O—(CH₂)_n—, —(CH₂)_n—S—S—(CH₂)_n—, —(CH₂)_n—, radical; B is a polymethylene radical having 3 to 10 carbon atoms and is optionally substituted by one or two alkyl substitutes having 1 to 12 carbon atoms, or is a xylylene radical, a —CH₂—CHOH—CH₂— radical or a —(CH₂)_n—O—(CH₂)_n— radical, n being 2 or 3, and X is a chlorine, iodine or bromine atom.

4. A method according to claim 1, in which R is a methyl radical, R' is an alkyl radical having 1 to 16 carbon atoms, a benzyl radical or a cyclohexyl radical, or R and R' together represent the —(CH₂)₂—O—(CH₂)₂ radical, A is a xylylene radical or a polymethylene radical having 2 to 12 carbon atoms which is optionally substituted by one or two alkyl substituents having 1 to 12 carbon atoms; B is a polymethylene radical having 3 to 10 carbon atoms which is optionally substituted by one or two alkyl substituents having 1 to 12 carbon atoms, or a xylylene radical; and X is a chlorine, iodine or bromine atom.

40 5. A method according to claim 1, in which R=R'=CH₃, X=Br and A represents

and
$$B = -(CH_2)_3$$
; or

```
A = -(CH_2)_2 and B represents -(CH_2)_4, -(CH_2)_6, -(CH_2)_{10} or
                                                                        or A = -(CH_2)_s— and B represents -(CH_2)_s—, -(CH_2)_s, -(CH_2)_{10}— or
                                                                                                                                                                                                                                                                                                                          CH<sub>2</sub>--{
                                                                                    A = -(CH_2)_6— and B represents -(CH_2)_3—, -(CH_2)_6—, -(CH_2)_1—, -(CH_2)_2—-(CH_2)_3—
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       —(CH<sub>2</sub>)<sub>4</sub>—, —(CH<sub>2</sub>)<sub>5</sub>—,
                                                                                                                                                                                                                                                                                                                                                  CH,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ĊH.
                                                                     an o or p-xylylene radical; or
                                                                                                 A = -(CH_2)_s and B = -(CH_2)_s, or -(CH_2)_s; or A = -(CH_2)_{10} and B = -(CH_2)_4 or -(CH_2)_{10}.

6. A method according to claim 1, in which R=R'=CH<sub>8</sub>, X is an iodine atom,
  10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10
                                                                     A = -(CH_2)_0 and B = -(CH_2)_3
                                                                                                  7. A method according to claim 1, in which R and R' together represent the
                                                                 **Note that the containing to claim 1, in which R and R together represent the -(CH_2)_2— radical, X=Br, A represents -(CH_2)_4— or -(CH_2)_{10}— and B represents -(CH_2)_3—, or A = -(CH_2)_4— and B = -(CH_2)_4.

8. A method according to claim 1, in which R = R' = CH_3, X = CI, and A = -(CH_2)_4— and B = -(CH_2)_4— or -(CH_2)_2—. or A = -(CH_2)_3— and A = -(CH_2)_2—. or A = -(CH_2)_3— and A = -(CH_2)_2—.
     15
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                15
                                                                                    9. A method according to claim 1, in which R=R'=CH_3, X=Br, and A=(CH_2)_2-S-S-(CH_2)_2- and B represents -(CH_2)_3-, -(CH_2)_4-,                                                                                    (CH_2)_3, -(CH_2)_6, -(CH_2)_{10}, -(CH_2)_{10}, -(CH_2)_{20}, -(C
    20
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               20
                                                                 and B represents -(CH_2)_3—; or A represents -(CH_2)_2—O—(CH_2)_2— and B represents a p-xylylene radical, a -(CH_2)_2—O—(CH_2)_3— radical or a -(CH_2)_4—radical; or A = -(CH_2)_3—, -(CH_2)_4—, -(CH_2)_5—, -(CH_2)_6—, -(CH_2)_1— or p-xylylene and B = -(CH_2)_4—CHOH—-(CH_2)_4—rad B = -(CH_2)_5—Represents a p-xylylene radical; or A = -(CH_2)_3—or p-xylylene and B = -(CH_2)_4—rad B = -(CH_2)_5—rad B = -(CH_2)_5
    25
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              25
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                                                                10. A method according to claim 1, in which K = CH_3, K' = C_4H_4 and K = -(CH_2)_2 = -S_4 = -(CH_2)_2 = -CH_2 = -(CH_2)_2 = -(CH_2)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         35
                                                                 B= -(CH_2)_2-O-(CH_3)_2-.

11. A method according to claim 1, in which R= -CH_2-CH_2OH, R'=CH_3
                                                                  and A = -(CH_2)_3—and B = -(CH_2)_6—, -(CH_2)_{10}—or
                                                                                                                                                                                                                                                       --CH<sub>2</sub>--CHOH--CH<sub>2</sub>--; or
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         40
                                                             A=m-xylylene and B= -(CH_2)_6; or R'=C<sub>4</sub>H<sub>3</sub>, A= -(CH_2)_6— and B= -(CH_2)_2—O--(CH_2)_2—; or R'=C<sub>4</sub>H<sub>17</sub>, A= -(CH_2)_4— and B=p-xylylene or -CH_2—CHOH—CH<sub>2</sub>—; or R'=C<sub>12</sub>H<sub>25</sub>, A= -(CH_2)_6— and B= -(CH_2)_6—; or R and R' together represent the divalent radical -(CH_2)_2—O--(CH_2)_2— or -(CH_2)_6—, A= -(CH_2)_6— and B= -(CH_2)_6—CH<sub>2</sub>—CHOH—CH<sub>2</sub>—.
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                                                              12. A method according to claim 1, in which either R=R'=CH_3, X=Br and A=-(CH_2)_4 and B=-(CH_2)_7, -(CH_2)_8, or o-xylylenyl, or A=-(CH_2)_3 and B=-(CH_2)_2.
                                                                or CH<sub>3</sub> A=
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     50
                                                           and B = -(CH_2)_z—O—(CH_1)_z— or —CH<sub>2</sub>—CHOH—CH<sub>2</sub>—, or A = p-xylylidenyl and B = -CH_2—CHOH—CH<sub>2</sub>—.
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 $-(CH_2)_3$ and $B = -(CH_2)_4$, or $A = -(CH_2)_6$ and B =

13. A method according to claim 1, in which R=CH₃, R'=n-propyl, X=Br and

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—(CH₂)₃—CH—, CH₃ o-xylylenyl or $-(CH_2)_4$, or A=m-xylylenyl and $B=-(CH_2)_{10}$.

14. A method according to claim 1, in which $R=CH_3$, R'= isopropyl, X=Br, $A=-(CH_2)_3$ — and $B=-(CH_2)_3$ — or $-(CH_2)_7$ —.

15. A method according to claim 1, in which $R=CH_3$, R'= benzyl, $B=-CH_2$ —CHOH—CH₂—, and $A=-(CH_2)_3$ — if X=Br and $A=-(CH_2)_6$ — if 5 5 X=CI. 10 16. A method according to claim 1, in which $R=CH_3$, $R'=C_4H_7$, X=Br, A=m-xylylenyl and $B=-CH_2-CHOH-CH_2-$. 10 17. A method according to claim 1, in which R and R' together represent a —(OH₂),— group, X=Br, B represents —CH₂—CHOH—CH₂— and A represents 15 or m-xylylenyl. 15 18. A method according to claim 2 in which R is a C₁ to C₅ alkyl group. 19. A method according to any one of the preceding claims in which the polymer is applied in the form of an aqueous, alcoholic or aqueous-alcoholic solution, cream, gel or emulsion, or an aerosol containing a propellant. 20. A method according to any one of the preceding claims, in which the polymer 20 20 is applied in the form of a composition containing from 0.5 to 10% by weight of polymer. 21. A method according to claim 19 or 20 in which the polymer is as defined in claim 4 or 18.

22. A method according to any one of the preceding claims in which the polymer 25 25 is applied to the hair before an anionic and/or non-ionic shampoo or before an oxidation dyeing process followed by an anionic and/or non-ionic shampoo. 23. A composition suitable for application to the hair and/or skin which comprises at least one quaternised polymer as defined in any one of claims 1 to 18 and at least 30 one cosmetic adjuvant which is: 30 (i) a cosmetic resin such that the composition is in the form of a hair lacquer or wave-setting lotion; (ii) a hair dye; (iii) a hair restructuring agent; 35 (iv) a surface-active agent such that the composition is in the form of a 35 shampoo; or (v) an oil such that the composition is in the form of an emulsion for application to the skin. 24. A composition according to claim 23 which is in the form of an aqueous or 40 alcoholic solution containing a cosmetic resin. 40 25. A composition according to claim 23 which contains a cosmetic resin and an aerosol propellant. 26. A composition according to claim 23 which is in the form of a cream and contains a hair dye.

27. A composition according to claim 23 which contains an oil and is a cream, a sun-cream, a tinting cream or a cleansing milk.
28. A composition according to any one of claims 23 to 27 in which the polymer

is present in an amount from 0.5 to 10% by weight.

29. A composition according to any one of claims 23 to 28 which contains at

least one perfume, preservative, sequestering agent or thickener.

30. A composition according to any one of claims 23 to 29 in which the polymer

is one specifically identified herein.

31. A composition according to claim 30 in which the polymer is one defined in

claim 4 or 18.

55 32. A composition according to claim 23 substantially as hereinbefore described.

33. A method according to any one of claims 1 to 22 which comprises applying a composition as claimed in any one of claims 23 to 32.

34. A method according to claim 1 substantially as hereinbefore described.

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Printed for Her Majesty's Stationery Office by the Courier Press, Leamington Spa, 1978.

Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.